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(54) THE: NEW PEPTIDE DERIVATIVES

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The invention relates to new comp-inhibitors of trypath-like scripe protesses synthesis, pharmacoutical compositions or BEST AVAILABLE COPY

(51) International Patent

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New peptide derivatives

This invention relates to new competitive inhibitors of trypsin-like serine proteases, especially thrombin and kininogenases such as kallikrein, their synthesis, pharmaceutical compositions containing the compounds as active ingredients, and the use of the compounds as thrombin inhibitors and anticoagulants and as antiinflammatory inhibitors, respectively.

The invention also relates to novel use of compounds as starting materials in synthesis of a serine protease inhibitor. Furthermore the invention relates to a novel structural fragments in serine protease inhibitors.

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#### BACKGROUND

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Blood coagulation is the key process involved in both haemostasis (i.e. prevention of blood loss from a damaged vessel) and thrombosis (i.e. the pathological occlusion of a blood vessel by a blood clot). Coagulation is the result of a complex series of enzymatic reactions, where one of the final steps is conversion of the proenzyme prothrombin to the active enzyme thrombin.

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Thrombin plays a central role in coagulation. It activates platelets, it converts fibrinogen into fibrin monomers, which polymerise spontaneously into filaments, and it activates factor XIII, which in turn crosslinks the polymer to insoluble fibrin. Thrombin further activates factor V and factor VIII in a positive feedback reaction. Inhibitors of thrombin are therefore expected to be effective anticoagulants by inhibition of platelets, fibrin formation and fibrin

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stabilization. By inhibiting the positive feedback

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mechanism they are expected to excert inhibition early in the chain of events leading to coagulation and thrombosis.

Kininogenasas are serine proteases that act on kininogens to produce kinins (bradykinin, kallidin, and Met-Lys-bradykinin). Plasma kallikrein, tissue kallikrein, and mast cell tryptase represent important kininogenases.

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30 25 20 15 events in asthma, rhinitis, and intestinal diseases. contribute to kinin formation and other pathogenic al., Am. Rev. Respir. Dis., 1992, 146:1535-1542) to mast cell tryptase will be released (Salomonsson et and inflammatory bowel diseases. Particulary in allergy many diseases including asthma, rhinitis, common cold, 47:993-1000). Plasma exudation is thus a feature of factors (Persson et al., Editorial, Thorax, 1992, inflammation, whether it is allergy, infection or other of the mechanisms that are involved in the process is ongoing. Plasma exudation occurs independent continually as long as the active plasma exudation plasma-derived kininogens inevitably will be all the protein systems of circulating blood. The into the tissue. The ensuing plasma exudate contains the blood vessels resulting in extravasation of plasma process is associated with increased permeability of Kinins (bradykinin, kallidin) are generally involved in interacting with different kallikreins, forming kinins inflammation. For example, the active inflammation

The kinins are biologically highly active substances with smooth muscle effects, sectretory effects, neurogenic effects, and actions that may perpatuate inflammatory processes including activation of phospholipase A<sub>2</sub> and increasing vascular permeability. The latter action potentially induces a vicious circle

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with kinins providing for the generation of more kinins

Tissue kallikrein cleaves primarily low molecular weight kininogen to produce kallidin and plasma kallikrein preferably releases bradykinin from high molecular weight kininogen.

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#### PRIOR ART

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Inhibitors of thrombin based on the amino acid sequence around the cleavage site for the fibrinogen Aa chain were first reported by Blombäck et al. in J. Clin. Lab. Invest. 24, suppl 107, 59, (1969), who suggested the sequence Phe-Val-Arg (P9-P2-P1, herein referred to as the P3-P2-P1 sequence) to be the best inhibitor.

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In US 4,346;078 has 5. Bajusz et al. described the thrombin inhibitor H-DPhe-Pro-Agm, a dipeptidyl derivative vith an aminoalkyl guanidine in the Plposition.

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Inhibitors of thrombin based on peptide derivatives with a cyclic aminoalkyl quanidine, e.g. 3-aminomethyl-1-amidinopiperidine, in the P1-position have been disclosed in EP-A2-0,468,231.

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In EP-A2-0,185,390 has S. Bajusz et. al. disclosed that replacing the agmatine with an arginine aldehyde gave a thrombin inhibitor which had much higher potency.

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Inhibitors of kallikrein based on the amino acid sequence around the cleavage site Arg-Ser have been reported earlier.

The arginine chloromethyl ketones H-DPro-Phe-Arg-CH<sub>2</sub>Cl

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and H-D Phe-Phe-Arg-CH<sub>2</sub>Cl were reported as plasma kallikrein inhibitors by Kettner and Shaw in Biochemistry 1978, 17:4778-4784 and Meth. Enzym. 1981, 80:826-842.

Likewise, esters and amides containing the H-DPro-Pha-Arg sequencs were reported by Pareed et al. in Ann. N.Y. Acad. Sci. 1981, 370:765-784 to be plasma kallikrein inhibitors.

Inhibitors of serine protesses that are based on electrophilic ketones instead of aldehydes in the Piposition are described in the following patent documents:

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EP-A2-0,195,212 describing peptidyl a-keto esters and amides, EP-A1-0,162,002 describing fluoroalkylamide ketones and EP-A2-0,164,144 describing  $\alpha,\beta,\delta$ - triketo compounds possessing different peptidase inhibiting properties.

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Inhibitors of trypsin-like serine protesses, such as thrombin and kallikrein, based on C-terminal boronic acid derivatives of arginine and isothiouronium analogues thereof have been revealed in EP-A2-

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WO 92/04371 describing kininogenase inhibitors, e.g. kallikrein inhibitors based on derivatives of arginine.

EP-A1-0,530,167 describing α-alkoxy ketone derivatives of arginine as thrombin inhibitors.

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### 35 DISCLOSURE OF THE INVENTION

An object of the present invention is to provide novel

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rectally, topically e.g. dermally, or via the kininogenase inhibitors which can be given orally, inhalation route. further object of the invention is to obtain rhinitis, urticaria, inflammatory bowel disease, and inhibitors which are orally bicavailable and selective arthritis. A further object is to obtain thrombin treatment of inflammatory disorders e.g. asthma, disease, as well as inhibition of kininogenases for hypercoagulable states, e.g. following angioplasty and in inhibiting thrombin over other serine proteases. A thrombin is believed to play a role, e.g. Alzheimers coronary bypass operations, and other situations where thrombosis, general hypercoagulable states and local particular myocardial infarction and cerebral thrombosis, pulmonary embolism, arterial thrombosis, in treatment of thromboembolic diseases such as venous specifically anticoagulants for prophylaxis and their enzyme i.e. causing reversible inhibition. More compounds with competitive inhibitory activity towards especially anticoagulantia and antiinflammatory and potent trypsine-like serine protease inhibitors,

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serine proteases, especially thrombin and kininogenases such as kallikrein: compounds of the general Formula I, either as such or including stereoisomers, are potent inhibitors of in the form of physiologically acceptable salts, and According to the invention it has been found that

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wherein:

Al represents a structural fragment of Formula IIa, IIb, IIc, IId or IIe;

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25 wherein:

k is an integer 0, 1, 2, 3 or 4;

is an integer 1, 2, 3 or 4;

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q is an integer 0, 1, 2 or 3;

position which is alpha to the carbonyl group, and the alpha substituent is a group  ${
m R}^{17}-({
m CH}_2)_p^-$ , wherein p is 4 carbon atoms and is possibly substituted in the R1 represents H, an alkyl group having 1 to 4 carbon atoms, or  $\mathbb{R}^{11}$ 00C-alkyl-, where the alkyl group has 1 to

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0,1 or 2 and R<sup>17</sup> is methyl, phenyl, OH, COOR<sup>12</sup>, CONHR<sup>12</sup>, where R<sup>12</sup> is H or an alkyl group having 1 to 4 carbon atoms, and R<sup>11</sup> is H or an alkyl group having 1 to 6 carbon atoms, or

 $R^1$  represents  $Ph(4-COOR^{12})-CH_2^{-}$ , where  $R^{12}$  is as defined above, or

B

R<sup>1</sup> represents R<sup>13</sup>-NH-CO-alkyl-, where the alkyl group has 1 to 4 carbon atoms and is possibly substituted alpha to the carbonyl with an alkyl group having 1 to 4 carbon atoms and where R<sup>13</sup> is H or an alkyl group having 1 to 4 carbon atoms or -CH<sub>2</sub>COOR<sup>12</sup>, where R<sup>12</sup> is as defined above, or

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R represents R 1200C-CH 2-00C-alkyl-, where the alkyl group has 1 to 4 carbon atoms and is possibly substituted alpha to the carbonyl with an alkyl group having 1 to 4 carbon atoms and where R 12 is as defined above, or

 $\rm R^1$  represents  $\rm R^{14}So_3-$ ,  $\rm Ph(4-COOR^{12})-So_2-$ ,  $\rm Ph(3-COOR^{12})-So_2-$ , where  $\rm R^{12}$  is as defined above and  $\rm R^{14}$  is an alkyl group having 1-4 carbon atoms, or

 $R^1$  represents -CO- $R^{15},$  wherein  $R^{15}$  is an alkyl group having 1-4 carbon atoms, or

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R<sup>1</sup> represents -co-ogi<sup>5</sup>, where R<sup>15</sup> is as defined above,

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 $R^1$  represent -CO-  $(CH_2)_p$ -COOR  $^{12}$  , where  $R^{12}$  is as defined above and p is an interger 0, 1 or 2, or

35 R<sup>1</sup> represents -CH<sub>2</sub>PO(OR<sup>16</sup>)<sub>2</sub>, -CH<sub>2</sub>SO<sub>2</sub>H or -CH<sub>2</sub>-(5-(1H)-tetrazolyl), where R<sup>16</sup> is, individually at each occurrence, H, methyl or ethyl;

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R<sup>2</sup> represents H or an alkyl group having 1 to 4 carbon atoms or R<sup>21</sup>OOC-alkyl-, where the alkyl group has 1 to 4 carbon atoms and, where R<sup>21</sup> is H or an alkyl group having 1 to 4 carbon atoms;

R<sup>3</sup> represents an alkyl group having 1-4 carbon atoms, and the alkyl group may or may not carry one or more flourine atoms, or

10 R<sup>3</sup> represents a cyclopentyl, cyclohexyl- or a phenyl group which may or may not be substituted with an alkyl group having 1 to 4 carbon atoms, or

R<sup>3</sup> represents a phenyl group substituted with a OR<sup>31</sup>
15 group, where R<sup>31</sup> is H or an alkyl group heving 1 to 4
carbon atoms and k is 0, 1, or

 $R^3$  represents a 1-naphthyl or 2-naphthyl group and k is 0, 1, or

 $\mathbf{R}^2$  represent a cis- or trans-decalin group and k is 0, 1, or

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R<sup>3</sup> represents 4-pyridyl, 3-pyrrolidyl or 3-indolyl which may or may not be substituted with a OR<sup>31</sup> group, where R<sup>31</sup> is as defined above and k is 0, 1, or

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 $R^3$  represents Si(Me), or  $CH(R^{32})_2$  , wherein  $R^{32}$  is a cyclohexyl- or a phenyl group;

 $R^4$  represents H, an alkyl group having 1 to 4 carbon atoms, a cyclohexyl- or a phenyl group;

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A<sup>2</sup> represents a structural fragment of Pormula IIIe,

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wherein:

p is an interger 0, 1 or 2;

Y represents a methylene group, or m is an integer 1, 2, 3 or 4;

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or may or may not be unsaturated, or atoms, a hydroxy group or an oxo group in position 4, membered ring may or may not carry one or two fluorine Y represents an ethylene group and the resulting 5-

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heteroatom functionality in position 4, or Y represents -CH2-0-, -CH2-S-, -CH2-SO-, with the

S G

an alkyl group with 1 to 4 carbon atoms, or unsaturated in position 4 and 5, or carry in position 4 two fluorine atoms in one of positions 4 or 5 or be Y represents a n-propylene group and the resulting 6fluorine atom, a hydroxy group or an oxo group, carry membered ring may or may not carry in position 5 one

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Y represents -CH2-0-CH2-, -CH2-S-CH2-, -CH2-SO-CH2-, or

Y represent -CH2-CH2-CH2-CH2-;

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R<sup>3</sup> is as defined above;

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R<sup>5</sup> represents H or an alkyl group having 1 to 4 carbon atoms, or

R<sup>51</sup> is H or an alkyl group having 1 to 4 carbon atoms;  $R^5$  represents  $-(CH_2)_p$ -COOR<sup>51</sup>, where p is 0, 1 or 2 and

n is an integer 0, 1, 2, 3 or 4; av

5 IVe or IVe B represents a structural fragment of Formula IVa, IVb,

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wherein:

r is an interger 0 or 1;

25 X1 represent CH2, NH or is absent;

X2 represents CH2, NH or C=NH;

30 NH-C(NH)-NH2 or CH-CH2-C(NH)-NH2; x3 represents NH, C=NH, N-C(NH)-NH2, CH-C(NH)-NH2, CH-

X4 represents CH2 or NH;

Preferred combinations of  $X^1$ ,  $X^2$ ,  $X^3$ ,  $X^4$  and r are

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 $x^1$ ,  $x^2$  and  $x^4$  are CH<sub>2</sub>,  $x^3$  is CH-C(NH)-NH<sub>2</sub> and x is 0, 1,

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 $\chi^1$ ,  $\chi^2$  and  $\chi^4$  are CH2,  $\chi^3$  is N-C(NH)-NH2 and r is 0, 1,

 $x^1$  and  $x^3$  are NH,  $x^2$  is C=NH,  $x^4$  is CH2 and r is 0, 1,

 $\chi^{1}$  is  $\mathrm{CH}_{2}$  ,  $\chi^{2}$  and  $\chi^{4}$  are NH,  $\chi^{3}$  is C-NH and r is 1, or  $\chi^3$  and  $\chi^4$  are CH2,  $\chi^2$  is C=NH,  $\chi^3$  is NH and r is 0, 1,  $\chi^1,~\chi^2$  and  $\chi^4$  are  $CH_2,~\chi^3$  is  $CH-NH-C(NH)-NH_2$ 

 $x^1$ is absent,  $x^2$  and  $x^4$  are  $cH_2$ ,  $x^3$  is  $N-C(NH)-NH_2$  and r $\chi^1$  is absent,  $\chi^2$  and  $\chi^4$  are  $CH_2$ ,  $\chi^3$  is  $CH-C(NH)-NH_2$  and and r is 0, 1, or 9

 $\chi^1$  ,  $\chi^2$  and  $\chi^4$  are CH<sub>2</sub>,  $\chi^3$  is CH-C(NH)-NH<sub>2</sub> and r is 1,  $\chi^1$  ,  $\chi^2$  and  $\chi^4$  are CH<sub>2</sub>,  $\chi^3$  is N-C(NH)-NH<sub>2</sub> and r is 0 or Particularly preferred combinations of  $x^1,\ x^2,\ x^3,\ x^4$ and r are 13

 $x^{1}$  is absent,  $x^{2}$  and  $x^{4}$  are  $c_{\rm H_{2}}$ ,  $x^{3}$  is N-C(NH)-NH<sub>2</sub> and X1 and X3 are NH, X2 is C=NH, X4 is CH2 and r is 1; 20

 $\chi^5$  represents C(NH)-NH $_2$  or NH-C(NH)-NH $_2$ ;

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Re is H or an alkyl group having 1-4 carbon atoms;

X<sup>6</sup> represents CH or N;

- Compounds of Formula I having S-configuration on the  ${\tt A}^2$ amino acid are preferred ones, of those compounds also having R-configuration on the A<sup>1</sup> amino acid are particularly preferred ones. 30
- unless specified otherwise. An alkyl group having 1 to In the present context the term "an alkyl group having 1 to 4 carbon atoms" may be straight or branched 35

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4 carbon atoms may be methyl, ethyl, n-propyl, ipropyl, n-butyl, i-butyl, s-butyl and t-butyl.

n-butyl, i-butyl, s-butyl, t-butyl, n-pentyl, i-pentyl, In the present context the term "an alkyl group having 1 to 6 carbon atoms" may be straigh or branched unless carbon atoms may be methyl, ethyl, n-propyl, i-propyl, unsaturation is referred to, a carbon-carbon double specified otherwise. An alkyl group having 1 to 6 t-pentyl, neo-pentyl, n-hexyl or 1-hexyl. When

bond is intended.

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The wavy lines on the carbon atom in the carbonyl group IVb, IVc, IVd signfy the bond position of the fragment. in formulas IIa, IIb, IIc, IId, IIe, IIIa, IIIb, IIIc, on the carbon atom in the ring system in formulas IVa, on the nitrogen atom in formulas IIIa, IIIb, IIIc and

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Abbreviations are listed at the end of this specification.

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compounds of the general Formula Ia, either as such or in the form of physiologically acceptable salts, and including stereoisomers, are potent inhibitors of According to the invention it has been found that

thrombin:

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wherein:

A<sup>1</sup> represents a structural fragment of Formula IIa, IIb, IIc or IId, preferably IIa or IIb; 35

k is an integer 0, 1, 2, 3 or 4, preferably 0, 1;

q is an integer 0, 1, 2 or 3, preferably 1;

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carbon atoms, or atoms, and  $\mathbb{R}^{11}$  is H or an alkyl group having 1 to 6 where  $\mathbb{R}^{12}$  is H or an alkyl group having 1 to 4 carbon 0,1 or 2 and R<sup>17</sup> is methyl, phenyl, OH, COOR<sup>12</sup>, CONHR<sup>12</sup> position which is alpha to the carbonyl group, and the alpha substituent is a group R17-(CH2)p-, wherein p is carbon atoms and is possibly substituted in the atoms,  $R^{11}$ 00C-alkyl-, where the alkyl group has 1 to 4  $R^2$  represents H, an alkyl group having 1 to 4 carbon

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 $\mathbb{R}^1$  represents Ph(4-COOR $^{12}$ )-CH $_2$ -, where  $\mathbb{R}^{12}$  is as defined

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25 20 defined above, or 1 to 4 carbon atoms or "CH2COOR12 where R12 is as carbon atoms and where R<sup>13</sup> is H or an alkyl group having alpha to the carbonyl with an alkyl group having 1 to 4 has 1 to 4 carbon atoms and is possibly substituted  $\mathbb{R}^1$  represents  $\mathbb{R}^{11}$ -NH-CO-alkyl-, where the alkyl group

having 1 to 4 carbon atoms and where  $\mathbb{R}^{12}$  is as defined substituted alpha to the carbonyl with an alkyl group group has 1 to 4 carbon atoms and is possibly  ${
m R}^1$  represents  ${
m R}^{12}$ 00C-CH $_2$ -00C-alkyl-, where the alkyl

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 $^{\circ}$  and  $\mathbb{R}^{14}$  is an alkylgroup having 1-4 carbon atoms, or  $\mathrm{SO}_2-$ ,  $\mathrm{Ph}(2\mathrm{-COOR}^{12})\mathrm{-SO}_2-$  where  $\mathrm{R}^{12}$  is as defined above  $R^1$  represents  $R^{14}SO_2$ -,  $Ph(4-COOR^{12})-SO_2$ -,  $Ph(3-COOR^{12})$ -

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R<sup>1</sup> represents -co-R<sup>15</sup>, wherein R<sup>15</sup> is an alkyl group

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having 1-4 carbon atoms, or

R<sup>1</sup> represents -CO-OR<sup>15</sup>, where R<sup>15</sup> is as defined above,

above and p is an interger 0, 1 or 2, or  $\mathbb{R}^1$  represent -CO-(CH<sub>2</sub>)<sub>p</sub>-COOR<sup>12</sup>, where  $\mathbb{R}^{12}$  is as defined

5 each occurrence, H, methyl or ethyl;  $-CH_2-(5-(1H)-tetrazolyl)$ , where  $R^{16}$  is, individually at  $\mathbb{R}^1$  represents  $-CH_2PO(OR^{16})_2$ ,  $-CH_2SO_3H$  or

group has 1 to 4 carbon atoms and R11 is H. Preferably R<sup>1</sup> represents R<sup>11</sup>00C-alkyl-, where the alkyl

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to 4 carbon atoms; 4 carbon atoms and  $\mathbb{R}^{21}$  is H or an alkyl group having 1 atoms, or  $R^{2}$ 00C-alkyl-, where the alkyl group has 1 to R<sup>2</sup> represents H or an alkyl group having 1 to 4 carbon

fluorine atoms, or R<sup>1</sup> represents an alkyl group having 1-4 carbon atoms, and the alkyl group may or may not carry one or more

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25 group having 1 to 4 carbon atoms, or group which may or may not be substituted with an alkyl R3 represents a cyclopentyl, cyclohexyl- or a phenyl

ü is 0, 1, or R<sup>3</sup> represents a 1- naphthyl or 2-naphthyl group and k

1, or  ${f R}^{f J}$  represent a cis- or trans-decalin group and k is 0,

35  $R^3$  represents Si(Me)<sub>3</sub> or CH( $R^{32}$ )<sub>2</sub>, wherein  $R^{32}$  is a cyclohexyl- or phenyl group;

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atoms, a cyclohexyl or a phenyl group, preferably a R4 represents an alkyl group having 1 to 4 carbon cyclohexyl or a phenyl group; A<sup>2</sup> represents a structural fragment of Formula IIIa, IIIb or IIIc, preferably IIIa; S.

wherein:

p is an interger 0, 1 or 2; 2 m is an integer 1, 2, 3 or 4, preferably 2, 3;

Y represents a methylene group, or

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membered ring may or may not carry and or two fluorine atoms, a hydroxy group or an oxo group in position 4, Y represents an ethylene group and the resulting 5or may or may not be unsaturated, or

Y represents -CH2-O-, -CH2-S-, -CH2-SO-, with the heteroatom functionality in position 4, or

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unsaturated in position 4 and 5, or carry in position 4 Y represents a n-propylene group and the resulting 6fluorine atom, a hydroxy group or an oxo group, carry membered ring may or may not carry in position 5 one two fluorine atoms in one of positions 4 or 5 or be an alkyl group with 1 to 4 carbon atoms, or

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Y represents -CH2-O-CH2-, -CH3-S-CH2-, -CH2-SO-CH2-, or

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Y represent -CH2-CH2-CH2-;

R3 represents an alkyl group having 1-4 carbon atoms, 35

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R³ represents a S1(Me), group;

R<sup>5</sup> represents H or an alkyl group having 1 to 4 carbon atoms, preferably H or a methylgroup, or  $R^5$  represents -(CH<sub>2</sub>)  $_p$ -COOR<sup>51</sup>, where p is 0, 1 or 2 and R<sup>51</sup> is H or an alkyl group having 1 to 4 carbon atoms, preferably p is 0 and R<sup>51</sup> is H;

n is an integer 0, 1, 2, 3 or 4, preferably 1, 2, 3; 2

B represents a structural fragment of Formula IVa, IVD, IVC or IVd, preferably IVa or IVb

wherein:

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 $\chi^1$ ,  $\chi^2$ ,  $\chi^3$ ,  $\chi^4$ ,  $\chi^5$  and  $\chi^6$  are as defined above;

r is an integer 0 or 1;

R<sup>6</sup> is H or an alkyl group having 1-4 carbon atoms, preferably H; 2

preferred combinations of  $X^1,\ X^2,\ X^3,\ X^4$  and r are

 $\chi^1$ ,  $\chi^2$  and  $\chi^4$  are  $CH_2$ ,  $\chi^3$  is  $CH-C(NH)-NH_2$  and r is 0 or 1, or 23

ö  $\chi^1$ ,  $\chi^2$  and  $\chi^4$  are  $cH_2$ ,  $\chi^3$  is N-C(NH)-NH $_2$  and r is 0

1, or

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X1 and X3 are NH, X2 is C=NH, X4 is CH2 and r is 0 or

 $\chi^2$  and  $\chi^4$  are  $\text{CH}_2^2$ ,  $\chi^2$  is C-NH,  $\chi^3$  is NH and r is 0 or 1, 33

X1 is CH2, X2 and X4 are NH, X3 is C-NH and r is 1, or

 $x^1$ ,  $x^2$  and  $x^4$  are  $CH_2$ ,  $x^3$  is CH-NH-C(NH)-NH $_2$  and r=0 or 1, or

or  $x^1$  is absent,  $x^2$  and  $x^4$  are  $\text{CH}_2$ ,  $x^3$  is  $\text{CH-C(NH)-NH}_2$  and r is 0,

or  $x^1$  is absent,  $x^2$  and  $x^4$  are  $CH_2$ ,  $x^3$  is  $N-C(NH)-NH_2$  and r is 0;

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Particularly preferred combinations of  $\chi^1$ ,  $\chi^2$ ,  $\chi^3$ ,  $\chi^4$  and r are

15  $X^1$  is absent,  $X^2$  and  $X^4$  are  $CH_2$ ,  $X^3$  is  $N-C(NH)-NH_2$  and r is 0, or

 $x^{1}$ ,  $x^{2}$  and  $x^{4}$  are  $CH_{2}$ ,  $x^{3}$  is  $CH-C(NH)-NH_{2}$  and r=1, or

20  $x^1$ ,  $x^2$  and  $x^4$  are  $CH_2$ ,  $x^3$  is N-C(NH)-NH<sub>2</sub> and r = 0 or 1, or

 $x^1$  and  $x^3$  are NH,  $x^2$  is C=NH,  $x^4$  is CH<sub>2</sub> r is 1;

25  $X^{S}$  represents  $C(NH)-NH_{2}$  or  $NH-C(NH)-NH_{2}$ , preferably  $C(NH)-NH_{2}$ ;

X<sup>6</sup> represents CH or N;

30 According to a preferred embodiment the invention relates to compounds of Formula Ia,

herein;

35 A<sup>1</sup> represents a structural fragment of Formula IIa,

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wherein:

k is 0 or 1;

 $\mathbb{R}^1$  represents  $\mathbb{R}^{11}$ 00C-alkyl-, where the alkyl group has 1 to 4 carbon atoms, particularly methylene, ethylene and  $\mathbb{R}^{11}$  is  $H_i$ 

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R2 represents H;

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R<sup>3</sup> represents a cyclohexyl group;

 ${\bf A^2}$  represents a structural fragment of Pormula IIIa, wherein:

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Y represents a methylene group, an ethylene group, or a n-propylene group and the resulting 6-membered ring may or may not carry in position 4 an alkyl group with 1 to 4 carbon atoms, preferably Y represents methylene, ethylene;

R<sup>5</sup> represents H;

20

B represents a structural fragment of formula IVa

wherein:

25

 $x^1$  is absent,  $x^2$  and  $x^4$  are  $CH_2$ ,  $x^3$  is N-C(NH)-NH<sub>2</sub>, r is 0 and n is 1 or 2;

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 $x^1$ , and  $x^3$  are NH,  $x^2$  is C=NH,  $x^4$  is CH<sub>2</sub>, r is 1 and n is 2, or

 $\mathbf{x}^1$ ,  $\mathbf{x}^2$  and  $\mathbf{x}^4$  are  $\mathrm{CH}_2$ ,  $\mathbf{x}^3$  is  $\mathrm{CH-C(NH)-NH}_2$ , r is 1 and n is 1, or

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 $x^1$ ,  $x^2$  and  $x^4$  are  $CH_2$ ,  $x^3$  is  $N-C(NH)-NH_2$ , r is 0 or 1

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and n is 1 or 2, or

More particularly preferred are compounds wherein B represents a structural fragment fo formula IVb wherein:

 $X^5$  represents C(NH)-NH<sub>2</sub>, R<sup>6</sup> is H, and n = 1

Preferred compounds of the invention are:

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HOOC-CH2-CH2-(R) C91-Aze-Pab HOOC-CH2-CH2-(R) Cg1-Pro-Pab HOOC-CH2-(R) Cg1-Pro-Pab ноос-сн2-(R) сд1-Аге-РаБ

(HOOC-CH2)2-(R)Cg1-Pro-Pab H-(R)Cgl-Pic-Pab 12

НООС-СН<sub>2</sub>- (R, S) СН (СООН) - (R) Cha-Aze-Pab HOOC-CH2-(R, 8) CH (COOH) - (R) C91-P1C-Pab ноос-сн<sub>2</sub>-(R) cha-Aze-Pab H-(R)Cha-Aze-Pab 20

HOOC-CH2-(Rors) CH (COOH) - (R) Cha-Aze-Pab/a HOOC-CH2-(ROFS) CH(COOH)-(R) Cha-Aze-Pab/b HOOC-CH2-NH-CO-CH2-(R) Cha-Aze-Pab HOOC-CH2-CH2-(R) Cha-Aze-Pab

HOOC-CH2-(RorS) CH (COOH) - (R) Cha-Pro-Pab/b HOOC-CH2-(Rors) CH(COOH)-(R) Cha-Pro-Pab/a ROOC-CH2-CH2-(Me) (R) Cha-Pro-Pab HOOC-CH2-CH2-(R) Cha-Pro-Pab HOOC-CH2-(Me) (R) Cha-Pro-Pab HOOC-CH2-(R) Cha-Pro-Pab H-(R)Cha-Pro-Pab 8 **5**2

HOOC-CH2-(Rors) CH(COOH)-(R) Cha-Pic-Pab/a HOOC-CH2-NH-CO-CH2-(R) Cha-Pro-Pab Stooc-cH2-cH2-CH2-(R) Cha-Pro-Pab Ph (4-COOH) -SO<sub>2</sub>-(R) Cha-Pro-Pab HOOC-CH2-(R) Cha-Pic-Pab H-(R) Cha-Pic-Pab 35

HOOC-CH2-(Rors) CH (COOH)-(R) Cha-Pic-Pab/b Me-00C-CH3-CO-(R) Cha-Pic-Pab HOOC-CH2-CH2-(R) Cha-Pic-Pab HOOC-CH2-CO-(R) Cha-Pic-Pab HOOC-CO-(R) Cha-Pic-Pab

H2N-CO-CH2-(R)Cha-Pic-Pab Ma-SO2-(R) Cha-Pic-Pab Boc-(R) Cha-Pic-Pab Ac-(R) Cha-Pic-Pab

HOOC-CH2-CH2-(R) Cha-(R, S) betaPic-Pab HOOC-CH2-CH2-(R) HOC-Aze-Pab HOOC-CH2-CH2-(R) Cha-Val-Pab H-(R) Cha-(R, S) betaPic-Pab HOOC-CH2-(R) Cha-Val-Pab H-(R) Hoc-Aze-Pab 15 ទ

HOOC-CH2-(R, S) CH (COOH) - (R) HOC-Pro-Pab HOOC-CH2-CH2-(R) Pro(3-(S) Ph) -Pro-Pab HOOC-CH2-(R) Pro(3-(S) Ph) -Pro-Pab HOOC-CH2-CH2-(R)Tic-Pro-Pab (HOOC-CH2) 2-(R) Hoc-Pic-Pab HOOC-CH2-(R) Hoc-Pic-Pab 20

HOOC-CH2-CH2-(R)Cgl-Aze-Pig

HOOC-CH2-(R) Cgl-Pro-Pig

HOOC- (R, S) CH (Me) - (R) Cha-Pro-Pab HOOC-CH2-(R) Cg1-Aze-Pac H-(R) Cg1-Aze-Pab H-(R) Cha-Aze-Pig H-(R) Cha-Pro-Pac H-(R)Cg1-11e-Pab 25

"HexOCC-CH2-(R) Cg1-Aze-Pab "Buccc-CH2-(R) Cg1-Aze-Pab Meooc-CH2-(R) Cg1-Aze-Pab Etooc-CH2-(R) Cg1-Aze-Pab H-(R)Cgl-Pro-Pac 30

HOOC-CH2-CH2-(R) CG1-Pro-Pac HOOC-CH2-CH2-(R) Cha-Aze-Pac HOOC-CH2-(R) Cha-Pro-Pac 33

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 $HOOC-CH_2-(R)$  Cha-Pic-(R,S) Itp H-(R)Cha-Pic-(R,S)Itp HOOC-CH<sub>2</sub>-(R) Cha-Aze-(R, 8) Itp HOOC-CH2-CH2(HOOC-CH2)-(R)Cha-Pro-Pig HOOC-CH2-(R)Cgl-Aze-(R,S)Itp

15 5 HOOC-CH2-(R) Cha-Pro-Rig HOOC-CH2-(R) Cgl-Aze-Rig H-(R)Cha-Pro-(R,S)Hig H-(R)Cgl-Pro-(R,S)Hig H-(R)Cgl-Aze-Rig HOOC-CH2-(R)Cgl-Pro-(R,S)Hig

20 H-(R)Cha-Aze-Dig H-(R)Cha-Pro-Dig H-(R)Cha-Pro-Mig H-(R)Cha-Pro-(R,S)Nig HOOC-CH2-(R) Cha-Pro-(8) Itp HOOC-CH2-CH2-(R) Cha-Aze-Rig

At present the particularly preferred compounds of formula Ia is

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30 HOOC-CH2-(R) Cha-Pro-Pig HOOC-CH2-(R) Cha-Pro-Pac EtOOC-CH2-(R)Cgl-Aze-Pab HOOC-CH2-(R) Cgl-Pro-Pig HOOC-CH2-(R) Cha-Pic-Pab HOOC-CH2-CH2-(R) Cha-Pro-Pab HOOC-CH2-(R) Cha-Pro-Pab HOOC-CH2-CH2-(R) Cha-Aze-Pab HOOC-CH2-(R)Cgl-Aze-Pab

refer to a substantially pure stereoisomer at the In the above tables of compounds, the letters /a and /b 35

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experimental part herein. "R,S" refers to a mixture of sterecisomers. identified for each compound with reference to the carbon atom noted "Rors". The stereoisomer can be

kininogenases: in the form of physiologically acceptable salts, and compounds of the general Formula Ib, either as such or including stereoisomers, are potent inhibitors of According to the invention it has been found that

 $-A^2$  NH-(CH<sub>2</sub>) -B

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wherein:

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or IIe, preferably IIa or IIb; A represents a structural fragment of formula IIa, IIb

20 wherein:

k is an integer 0, 1, 2, 3 or 4, preferably 0, 1;

q is an integer 0, 1, 2, or 3, preferably 1;

ä 25 0, 1 or 2 and  $\mathbb{R}^{17}$  is methyl, phenyl, OH, COOR<sup>12</sup>, carbon atoms, and  $\mathbb{R}^{11}$  is H or an alkyl group having 1 to 6 carbon atoms, or CONHR<sup>12</sup>, where  $\mathbb{R}^{12}$  is H or an alkyl group having 1 to 4 alpha substituent is a group  $R^{12}$ -(CH<sub>2</sub>)<sub>p</sub>-, wherein p is position which is alpha to the carbonyl group, and the 4 carbon atoms and is possibly substituted in the atoms, or R1100C-alky1-, where the alky1 group has 1 to R1 represents H, an alkyl group having 1 to 4 carbon

ü  $\mathbb{R}^1$  represents Ph(4-COOR<sup>12</sup>)-CH<sub>2</sub>-, where  $\mathbb{R}^{12}$  is H or an alkyl group having 1 to 4 carbon atoms, or

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carbon atoms and where R<sup>13</sup> is H or an alkyl group having alpha to the carbonyl with an alkyl group having 1 to 4  $R^1$  represents  $R^{13}$ -NH-CO-alkyl-, where the alkyl group has 1 to 4 carbon atoms and is possibly substituted 1 to 4 carbon atoms or -CH2COOR<sup>12</sup> where R<sup>12</sup> is as defined above, or

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having 1 to 4 carbon atoms and where  $R^{12}$  is as defined substituted alpha to the carbonyl with an alkyl group  $R^1$  represents  $R^{12}00C-CH_2-00C-alkyl-$ , where the alkyl group has 1 to 4 carbon atoms and is possibly above, or

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 $m R^1$  represents  $m R^{14} So_2^-$ ,  $m Ph(4-COOR^{12})$  - $m So_2^-$ ,  $m Ph(3-COOR^{12})$  - $SO_2$ ,  $Ph(2-COOR^{12})-SO_2$ -, where  $R^{12}$  is as defined above and R<sup>14</sup> is an alkylgroup having 1-4 carbon atoms, or 15

 $\mathbf{R}^1$  represents -co- $\mathbf{R}^{15}$ , wherein  $\mathbf{R}^{15}$  is an alkyl group having 1-4 carbon atoms, or

 $\mathbb{R}^1$  represents -CO-OR $^{15}$ , where  $\mathbb{R}^{15}$  is as defined above, 20

 $\mathrm{R}^1$  represent -co-(CH<sub>2</sub>) $_\mathrm{p}$ -COOR $^{12}$ , where  $\mathrm{R}^{12}$  is as defined above and p is 0, 1 or 2, or 25

-CH<sub>2</sub>-(5-(1H)-tetrazoly1), where  $R^{16}$  is, individually at R1 represents -CH2PO(OR16)2, -CH2SO3H or each occurrence, H, methyl or ethyl;

atoms or  $\mathbb{R}^{21}$ 00C-alkyl-, where the alkyl group has 1 to 4 carbon atoms and  $\mathbb{R}^{21}$  is H or an alkyl group having 1 to R2 represents H or an alkyl group having 1 to 4 carbon 4 carbon atoms;

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R<sup>3</sup> represents an alkyl group having 1-4 carbon atoms, and the alkyl group may or may not carry one or more

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fluorine atoms, or

group which may or may not be substituted with an alkyl R3 represents a cyclopentyl, cyclohexyl- or a phenyl

group having 1 to 4 carbon atoms, or

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R<sup>3</sup> represents a phenyl group substituted with a OR<sup>31</sup> group, where R<sup>11</sup> is H or an alkyl group having 1 to carbon atoms and k is 0, 1, or

 $R^3$  represents a 1-naphthyl or 1-naphthyl group and k is 0, 1, or ខ្ព

 $\mathbf{R}^3$  represent a cis- or trans-decalin group and k is

0,1, or

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which may or may not be substituted with a  $oR^{13}$  group, R³ represents 4-pyridyl, 3-pyrrolidyl or 3-indolyl where R31 is as defined above and k is 0, 1, or

 $R^3$  represents Si(Me), or CH(R32),, wherein  $R^{32}$  is a cyclohexyl- or phenyl group;

20

atoms, a cyclohexyl or a phenyl group, preferably H; R<sup>4</sup> represents H, an alkyl group having 1 to carbon

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 $\mathsf{A}^2$  represents a structural fragment of formula IIIb or IIIc, preferably IIIb

wherein: 30

m is an integer 1, 2, 3, or 4, preferably 2, 3; p is an integer 0, 1 or 2; R3 is as defined above;

n is an integer 0, 1, 2, 3 or 4, preferably 1,2,3; 39

 $x^1$ ,  $x^2$ ,  $x^3$ ,  $x^4$  are as defined above;

preferably H or a methyl group; R<sup>6</sup> is H or an alkyl group having 1-4 carbon atoms,

r is an integer 0 or 1;

10

preferred combinations of  $x^1$ ,  $x^2$ ,  $x^3$  and  $x^4$  are

 $x^1$ ,  $x^2$  and  $x^4$  are  $CH_2$ ,  $x^3$  is  $CH-C(NH)-NH_2$  and r is 0 or 1, or

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1, or x1, x2 and  $x^4$  are  $CH_2$ ,  $x^3$  is N-C(NH)-NH<sub>2</sub> and r is 0 or

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õ  $x^1$  and  $x^3$  are NH,  $x^2$  is C=NH,  $x^4$  is CH<sub>2</sub> and r is 0 or 1,

e K2 and X4 are CH2, X2 is C-NH, X3 is NH and r is 0 or 1,

25

 $x^1$  is  $CH_2$ ,  $x^2$  and  $x^4$  are NH,  $x^3$  is C=NH and r is 1, or

or 1, or  $x^2$  is absent,  $x^2$  and  $x^4$  are  $CH_2$ ,  $x^3$  is  $CH-C(NH)-NH_2$  $x^1$ ,  $x^2$  and  $x^4$  are  $CH_2$ ,  $x^3$  is  $CH-NH-C(NH)-NH_2$  and r is 0

30

is 0;  $x^1$  is absent,  $x^2$  and  $x^4$  are  $cH_2$ ,  $x^3$  is N-C(NH)-NH<sub>2</sub> and r

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and r is 0, or

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particularly preferred combinations of  $X^1$ ,  $X^2$ ,  $X^3$  and  $X^4$ 

유  $x^1$ ,  $x^2$  and  $x^4$  are  $CH_2$ ,  $x^3$  is  $CH-C(NH)-NH_2$  and r is 1

 $x^{1}$ ,  $x^{2}$  and  $x^{4}$  are  $CH_{2}$ ,  $x^{3}$  is N-C(NH)-NH<sub>2</sub> and r is 1;

X<sup>5</sup> represents C(NH)-NH<sub>2</sub> or NH-C(NH)-NH<sub>2</sub>, preferably

X<sup>6</sup> represents CH or N.

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C(NH)-NH2;

Preferred compound of the invention are:

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H-(R)Cha-Cha-Pab HOOC-CH2-(R)Phe-Cha-Pab H-(R)Phe-Cha-Pab HOOC-CH2-(R) Cha-Phe-Pab HOOC-CO-(R) Phe-Phe-Pab H-(R)Phe-Phe-Pab H-(R)Cha-Phe-Pab HOOC-CH2-(R)Phe-Phe-Pab HOOC-CH2-(R) Pro-Phe-Pab H-(R)Pro-Phe-Pab

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as kallikrein after oral or parenteral administration: proteases, especially thrombin and kininogenases such stereoisomers, are potent inhibitors of serine physiologically acceptable salts, and including general Formula V, either as such or in the form of Furthermore, it has been found that compounds of the

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HOOC-CH2-(R) Cha-Cha-Pab

 $A^1 - A^2 - NH - (CH_2)_B - B - D$ Formula V

wherein:

 $\mathtt{A}^1$  represents a structural fragment of Formula IIa, IIb, IIc, IId or IIe;

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wherein:

k is an integer 0, 1, 2, 3 or 4;

m is an integer 1, 2, 3 or 4;

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q is an integer 0, 1, 2 or 3;

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0,1 or 2 and  $R^{17}$  is  $\text{COOR}^{12},\ \text{CONHR}^{12},\ \text{where}\ R^{12}$  is H or an  $R^{\rm 1}$  represents  $R^{\rm 11}00C\text{--alkyl-,}$  where the alkyl group has 1 position which is alpha to the carbonyl group, and the alpha substituent is a group  $\mathbb{R}^{17}\text{-}(\text{CH}_2)_p\text{-},$  wherein p is to 4 carbon atoms and is possibly substituted in the group, and  $R^{1\,1}$  is H or an alkyl group having 1 to 6 alkyl group having 1 to 4 carbon atoms or a benzyl carbon atoms, or a benzyl group, or 35

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 $\mathrm{R}^1$  represents Ph(4-COOR $^{12}$ )-CH $_2$ -, where  $\mathrm{R}^{12}$  is as defined above, or

carbon atoms and where  $R^{13}$  is H or an alkyl group having alpha to the carbonyl with an alkyl group having 1 to 4  $R^{\rm l}$  represents  $R^{\rm l3-NH-CO-alkyl-},$  where the alkyl group has 1 to 4 carbon atoms and is possibly substituted 1 to 4 carbon atoms or  $-CH_2COOR^{12}$ , where  $R^{12}$  is as defined above, or

having 1 to 4 carbon atoms and where  $\mathbb{R}^{12}$  is as defined substituted alpha to the carbonyl with an alkyl group  $R^1$  represents  $R^{12}00C$ - $CH_2$ -00C-alkyl-, where the alkyl group has 1 to 4 carbon atoms and is possibly above, or

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R<sup>1</sup> represents R<sup>14</sup>SO<sub>2</sub>-, Ph(4-COOR<sup>12</sup>)-SO<sub>2</sub>-, Ph(3-COOR<sup>12</sup>)-SO2-, Ph(2-COOR12)-SO2-, where R12 is as defined above and  $\mathbb{R}^{14}$  is an alkyl group having 1-4 carbon atoms, or

 $R^{1}$  represents -CO- $R^{15},$  wherein  $R^{15}$  is an alkyl group having 1-4 carbon atoms, or 20

 $R^1$  represents -CO-OR $^{15}$ , where  $R^{15}$  is as defined above,

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 $R^1$  represent -CO-(CH,)  $_p\text{-COOR}^{12}$  , where  $R^{12}$  is as defined above and p is an interger 0, 1 or 2, or

atoms or  $R^{21}00C$ -alkyl-, where the alkyl group has 1 to 4 carbon atoms and, where  $R^{2\,1}$  is H, an alkyl group having  $R^2$  represents H or an alkyl group having 1 to 4 carbon 1 to 4 carbon atoms or a benzyl group; 30

R³ represents an alkyl group having 1-4 carbon atoms, and the alkyl group may or may not carry one or more flourine atoms, or 35

R<sup>3</sup> represents a cyclopentyl, cyclohexyl- or a phenyl group which may or may not be substituted with an alkyl group having 1 to 4 carbon atoms, or

 $R^3$  represents a phenyl group substituted with a  $OR^{31}$  group, where  $R^{31}$  is H or an alkyl group having 1 to 4 carbon atoms and k is 0, 1, or

 $\ensuremath{R^3}$  represents a 1-naphthyl or 2-naphthyl group and  $\ensuremath{\kappa}$  is 0, 1, or

 $\mathbf{R}^3$  represent a cis- or trans-decalin group and k is 0, 1, or

15  $R^3$  represents 4-pyridyl, 3-pyrrolidyl or 3-indolyl which may or may not be substituted with a  $OR^{31}$  group, where  $R^{31}$  is as defined above and k is 0, 1, or

 $R^3$  represents Si(He)<sub>3</sub> or CH( $R^{32}$ )<sub>2</sub>, wherein  $R^{32}$  is a cyclohexyl- or a phenyl group;

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 $R^4$  represents H, an alkyl group having 1 to 4 carbon atoms, a cyclohexyl- or a phenyl group;

25 A<sup>2</sup>, B and n are defined as described under Formula I above;

D is Z or (Z)<sub>2</sub>, wherein Z represents a benzyloxycarbonyl group.

The benzyloxycarbonyl group (Z or (Z)<sub>2</sub>) will bind to the amidino- or guanidino nitrogens present in B.

Preferred and particularly preferred combinations are the same as described for Formula I above.

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Furthermore, it has been found that compounds of the

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general Formula Va, either as such or in the form of physiologically acceptable salts, and including stereoisomers, are potent inhibitors of thrombin after oral or parenteral administration:

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 $A^1 - A^2 - NH - (CH_2)_n - B - D$ 

Formula Va

wherein:

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A<sup>l</sup> represents a structural fragment of Formula IIa, IIb, IIc or IId, preferably IIa or IIb;

wherein:

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k is an integer 0, 1, 2, 3 or 4, preferably 0, 1;

q is an integer 0, 1, 2 or 3, preferably 1;

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R<sup>1</sup> represents R<sup>11</sup>OOC-alkyl-, where the alkyl group has 1 to 4 carbon atoms and is possibly substituted in the position which is alpha to the carbonyl group, and the alpha substituent is a group R<sup>17</sup>-(CH<sub>2</sub>)<sub>p</sub>-, wherein p is 0,1 or 2 and R<sup>17</sup> is COOR<sup>12</sup>, CONHR<sup>12</sup>, where R<sup>12</sup> is H, an alkyl group having 1 to 4 carbon atoms or a benzyl group, and R<sup>11</sup> is H or an alkyl group having 1 to 6 carbon atoms, or a benzyl group, or

 $\mathbb{R}^1$  represents  $\mathbb{P}h(4\text{-}COO\mathbb{R}^{12})\text{-}CH_2\text{--}$ , where  $\mathbb{R}^{12}$  is as defined above, or

R<sup>1</sup> represents R<sup>13</sup>-NH-CO-alkyl-, where the alkyl group has 1 to 4 carbon atoms and is possibly substituted alpha to the carbonyl with an alkyl group having 1 to 4 carbon atoms and where R<sup>13</sup> is H or an alkyl group having

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 $R^{\rm 3}$  represents a 1- naphthyl or 2-naphthyl group and kgroup having 1 to 4 carbon atoms, or

is 0, 1, or

 $R^3$  represent a cis- or trans-decalin group and k is 0,

1, or

 $R^3$  represents  $S1(Me)_3$  or  $CH(R^{32})_2, \ \mbox{wherein} \ R^{32}$  is cyclohexyl- or phenyl group; ខ្ព

atoms, a cyclohexyl or a phenyl group, preferably a  $R^4$  represents an alkyl group having 1 to 4 carbon cyclohexyl or a phenyl group;  $\mathtt{A}^2$ , B and n are defined as described under Formula Ia above;

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D is 2 or (2)2;

20

Z represents a benzyloxycarbonyl group.

described for Formula Ia above but  $\mathbb{R}^{11}$  is H, an alkyl particularly preferred combinations are the same as group having 1 to 6 carbon atoms or a benzyl group. Preferred integers, groups or combinations and

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Preferred compounds having Formula Va are:

Bnooc-CH2-CH2-(R) Cg1-Pro-Pab(Z) Bnooc-CH2-CH2-(R) Cgl-Aze-Pab(Z) Bnooc-CH2-(R)Cgl-Pro-Pab(Z) (Bnooc-CH<sub>2</sub>)<sub>2</sub>-(R)Cgl-Pro-Pab(Z) Bnooc-CH2-(R)Cgl-Aze-Pab(Z) 32 30

Bnooc-CH<sub>2</sub>-(R,S)CH(COOBn)-(R)Cg1-Pic-Pab(Z) Bnooc-CH<sub>2</sub>-(R)Cha-Aze-Pab(Z)

 $m R^1$  represents  $m R^{14}So_2^-$ ,  $m Ph(4-cooR^{12})-So_2^-$ ,  $m Ph(3-cooR^{12}) SO_2^{-}$ , Ph(2-COOR<sup>12</sup>)-SO<sub>2</sub>- where R<sup>12</sup> is as defined above and  $\mathbb{R}^{14}$  is an alkylgroup having 1-4 carbon atoms, or

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having 1 to 4 carbon atoms and where  $\mathbb{R}^{12}$  is as defined

above, or

substituted alpha to the carbonyl with an alkyl group

 $R^1$  represents  $R^{12}00C$ - $CH_2$ -00C-alkyl-, where the alkyl

group has 1 to 4 carbon atoms and is possibly

1 to 4 carbon atoms or  $^{-CH_2COOR^{12}}$  where  $R^{12}$  is as

defined above, or

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 $R^1$  represents -CO- $R^{15}$ , wherein  $R^{15}$  is an alkyl group having 1-4 carbon atoms, or

 $R^1$  represents -CO-OR15, where  $R^{15}$  is as defined above,

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 $R^1$  represent -CO-(CH2)  $_p\text{-COOR}^{12}$  , where  $R^{12}$  is as defined 20

above and p is an interger 0, 1 or 2, or

preferably  $R^1$  represents  $R^{11}00C$ -alkyl-, where the alkyl group has 1 to 4 carbon atoms and  $\mathbb{R}^{11}$  is as defined

above. 25

atoms, or  $\mathbb{R}^{21}$ 00C-alkyl-, where the alkyl group has 1 to 4 carbon atoms and  $\mathbb{R}^{21}$  is H or an alkyl group having 1  $R^2$  represents H or an alkyl group having 1 to 4 carbon to 4 carbon atoms or a benzyl group;

30

 $R^3$  represents an alkyl group having 1-4 carbon atoms, and the alkyl group may or may not carry one or more fluorine atoms, or

group which may or may not be substituted with an alkyl  $\mathtt{R}^3$  represents a cyclopentyl, cyclohexyl- or a phenyl

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Bnooc-CH<sub>2</sub>-(R,S) CH(COOBn) - (R) Cha-Aze-Pab(Z)
Bnooc-CH<sub>2</sub>-(RorS) CH(COOBn) - (R) Cha-Aze-Pab(Z) /a
Bnooc-CH<sub>2</sub>-(RorS) CH(COOBn) - (R) Cha-Aze-Pab(Z) /b
Bnooc-CH<sub>2</sub>-CH<sub>2</sub>-(R) Cha-Aze-Pab(Z)
Bnooc-CH<sub>2</sub>-CH<sub>2</sub>-(R) Cha-Aze-Pab(Z)
Bnooc-CH<sub>2</sub>-(R) Cha-Pro-Pab(Z)
Bnooc-CH<sub>2</sub>-(Me) (R) Cha-Pro-Pab(Z)
Bnooc-CH<sub>2</sub>-(He) (R) Cha-Pro-Pab(Z)
Bnooc-CH<sub>2</sub>-CH<sub>2</sub>-(R) Cha-Pro-Pab(Z)
Bnooc-CH<sub>2</sub>-CH<sub>2</sub>-(R) Cha-Pro-Pab(Z)
Bnooc-CH<sub>2</sub>-CH<sub>2</sub>-(R) Cha-Pro-Pab(Z)
Bnooc-CH<sub>2</sub>-CH<sub>2</sub>-(R) Cha-Pro-Pab(Z)
Bnooc-CH<sub>2</sub>-(R,S) CH(COOBn) - (R) Cha-Pro-Pab(Z)
Bnooc-CH<sub>2</sub>-NH-CO-CH<sub>2</sub>-(R) Cha-Pro-Pab(Z)
Ph(4-COOH)-So<sub>2</sub>-(R) Cha-Pro-Pab(Z)

BnOOC-CH<sub>2</sub>-NH-CO-CH<sub>2</sub>-(R) Cha-Pro-Pab(Z)

Ph(4-COOH)-SO<sub>2</sub>-(R) Cha-Pro-Pab(Z)

Boc-(R) Cha-Pic-Pab(Z)

BnOOC-CH<sub>2</sub>-(R) Cha-Pic-Pab(Z)

BnOOC-CH<sub>2</sub>-(R, S) CH(COOBh) - (R) Cha-Pic-Pab(Z)

BnOOC-CH<sub>2</sub>-(CH) Cha-Pic-Pab(Z)

EtOOC-CO-(R) Cha-Pic-Pab(Z)

MeOOC-CH<sub>2</sub>-CO-(R) Cha-Pic-Pab(Z)

H<sub>2</sub>N-CO-CH<sub>2</sub>-(R) Cha-Pic-Pab(Z)

Ac-(R) Cha-Pic-Pab(Z)

Me-SO<sub>2</sub>-(R) Cha-Pic-Pab(Z)

BnOOC-CH<sub>2</sub>-(R) Cha-Val-Pab(Z)

BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R) Cha-(R,S) Val-Pab(Z)

BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R) Hoc-Aze-Pab(Z)

BnOOC-CH<sub>2</sub>-(R,S) CH(COOBn) - (R) Hoc-Pro-Pab(Z)

BnOOC-CH<sub>2</sub>-(R) Hoc-Pic-Pab(Z)

BnOOC-CH<sub>2</sub>-(R) Pro(3-(S) Ph) -Pro-Pab(Z)
BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R) Pro(3-(S) Ph) -Pro-Pab(Z)
BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R) Tic-Pro-Pab(Z)
BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R) Cg1-Aze-Pig(Z)<sub>2</sub>
BnOOC-CH<sub>2</sub>-(R) Cg1-Pro-Pig(Z)<sub>2</sub>

 $(BnOOC-CH_2)_2-(R)Hoc-Pic-Pab(Z)$ 

BnOOC-(R,S)CH(Me)-(R)Cha-Pro-Pab(Z)

MeOOC-CH<sub>2</sub>-(R)Cg1-Aze-Pab(Z)

EtOOC-CH<sub>2</sub>-(R)Cg1-Aze-Pab(Z)

 $BnOOC-CH_2-(R)Cgl-Aze-Pac(Z)$ 

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EtOOC-CH<sub>2</sub>-(R) Cgl-Aze-Pab(Z)

\*\*Buooc-CH<sub>2</sub>-(R) Cgl-Aze-Pab(Z)

5) CH (COOBn) - (R) Cha-Aze-Pah(Z)

nHexooc-CH2-(R) Cg1-Aze-Pab(Z)
Bnooc-CH2-(R) Cha-Pro-Pac(Z)
Bnooc-CH2-CH2-(R) Cg1-Pro-Pac(Z)
Bnooc-CH2-CH2-(R) Cha-Aze-Pac(Z)
Bnooc-CH2-CH2-(R) Cha-Aze-Pig(Z)
Bnooc-CH2-(R) Cha-Pro-Pig(Z)
Bnooc-CH2-(R) Cha-Pro-Pig(Z)
Bnooc-CH2-CH2-(R) Cha-Pro-Pig(Z)
Bnooc-CH2-CH2-(R) Cha-Pro-Pig(Z)
Bnooc-CH2-(R) Cha-Pro-Pig(Z)
Bnooc-CH2-(R) Cg1-Pro-Pig(Z)
Bnooc-CH2-(R) Cha-Pic-(R,S) Itp(Z)
Bnooc-CH2-(R) Cg1-Pro-(R,S) Hig(Z)
Bnooc-CH2-(R) Cg1-Aze-Rig(Z)
Bnooc-CH2-(R) Cg1-Aze-Rig(Z)

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Particularly preferred compounds are:

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 $BnOOC-CH_2-CH_2-(R)Cha-Aze-Rig(Z)$ 

BnOOC-CH<sub>2</sub>-(R) cgl-Aze-Pab(Z)
BnOOC-CH<sub>2</sub>-(R) cha-Pro-Pab(Z)
20 BnOOC-CH<sub>2</sub>-(R) cha-Pic-Pab(Z)
BnOOC-CH<sub>2</sub>-(R) Cgl-Pro-Pig(Z)<sub>2</sub>
EtoOC-CH<sub>2</sub>-(R) Cgl-Aze-Pab(Z)
BnOOC-CH<sub>2</sub>-(R) cha-Pro-Pac(Z)
BnOOC-CH<sub>2</sub>-(R) Cha-Pro-Pig(Z)<sub>2</sub>

Furthermore, it has been found that compounds of the general Formula Vb, either as such or in the form of physiologically acceptable salts, and including stereoisomers, are potent inhibitors of kallikrein after oral or parenteral administration:

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$$A^1 - A^2 - NH - (CH_2)_n - B - D$$

Formula Vb

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and  $\mathbb{R}^{14}$  is an alkylgroup having 1-4 carbon atoms, or

 $R^1$  represents -co- $R^{15}$ , wherein  $R^{15}$  is an alkyl group having 1-4 carbon atoms, or  $R^1$  represents -CO-OR $^{15}$ , where  $R^{15}$  is as defined above,

ör

 $R^1$  represent -co-(CH2)  $_p^{-\text{COOR}^{1\,2}},$  where  $R^{1\,2}$  is as defined above and p is 0, 1 or 2, or 9

atoms or  $\mathbb{R}^{21} \odot \mathbb{C} \mathbb{C}$ -alkyl-, where the alkyl group has 1 to 4 carbon atoms and  $\mathbb{R}^{21}$  is H, an alkyl group having 1 to 4  $\mathbb{R}^2$  represents H or an alkyl group having 1 to 4 carbon carbon atoms or a benzyl group;

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R<sup>3</sup> represents an alkyl group having 1-4 carbon atoms, and the alkyl group may or may not carry one or more fluorine atoms, or

group which may or may not be substituted with an alkyl  $R^3$  represents a cyclopentyl, cyclohexyl- or a phenyl group having 1 to 4 carbon atoms, or

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 $\mathrm{R}^3$  represents a phenyl group substituted with a  $\mathrm{OR}^{31}$ group, where  $\mathbb{R}^{31}$  is H or an alkyl group having 1 to carbon atoms and k is 0, 1, or 25

 $R^3$  represents a 1-naphthyl or 1-naphthyl group and k is

0, 1, or

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 $R^3$  represent a cis- or trans-decalin group and k is 0,1, or

which may or may not be substituted with a  $\mathtt{OR}^{31}$  group,  $R^3$  represents 4-pyridyl, 3-pyrrolidyl or 3-indolyl where  $\mathbb{R}^{31}$  is as defined above and k is 0, 1, or 35

q is an integer 0, 1, 2, or 3, preferably 1;

 $R^{\rm l}$  represents  $R^{\rm ll}00C\text{-alkyl-},$  where the alkyl group has 1 position which is alpha to the carbonyl group, and the to 4 carbon atoms and is possibly substituted in the

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alpha substituent is a group  $\mathrm{R}^{17}\text{-}(\mathrm{CH}_2)_{p^-},$  wherein p is 0, 1 or 2 and  $R^{17}$  is  $COOR^{12}$ ,  $CONHR^{12}$ , where  $R^{11}$  is H or

an alkyl group having 1 to 4 carbon atoms, and  $R^{11}$  is H or an alkyl group having 1 to 6 carbon atoms, or a benzyl group, or  $R^1$  represents  $Ph(4-COOR^{12})-CH_2-$ , where  $R^{12}$  is as defined above, or

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carbon atoms and where  $\mathbb{R}^{13}$  is H or an alkyl group having alpha to the carbonyl with an alkyl group having 1 to 4  $R^{1}$  represents  $R^{13}\text{-}NH\text{-}CO\text{-}alkyl\text{-},}$  where the alkyl group has 1 to 4 carbon atoms and is possibly substituted 1 to 4 carbon atoms or  $-CH_2COOR^{12}$  where  $R^{12}$  is as

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 $R^1$  represents  $R^{12} OOC - CH_2 - OOC - alkyl-, where the alkyl$ defined above, or 30

having 1 to 4 carbon atoms and where  $\mathbb{R}^{12}$  is as defined substituted alpha to the carbonyl with an alkyl group group has 1 to 4 carbon atoms and is possibly

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above, or

R<sup>1</sup> represents R<sup>14</sup>SO<sub>2</sub>-, Ph(4-COOR<sup>12</sup>)-SO<sub>2</sub>-, Ph(3-COOR<sup>12</sup>)-SO2, Ph(2-COOR12)-SO2-, where R12 is as defined above

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k is an integer 0, 1, 2, 3 or 4, preferably 0, 1;

wherein:

 $\mathtt{A}^1$  represents a structural fragment of formula IIa, IIb

or IIe, preferably IIa or IIb;

wherein:

 $\mathbb{R}^3$  represents  $\mathrm{Si}(\mathrm{Me})_3$  or  $\mathrm{CH}(\mathbb{R}^{32})_2$ , wherein  $\mathbb{R}^{32}$  is a cyclohexyl- or phenyl group;

atoms, a cyclohexyl or a phenyl group, preferably H;  $R^4$  represents H, an alkyl group having 1 to carbon

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above;  ${f A^2}$ ,  ${f B}$  and  ${f n}$  are defined as described under Formula Ib

D represents Z or (Z)2.

group having 1 to 6 carbon atoms or a benzyl group. described in Formula Ib above but R<sup>11</sup> is H, an alkyl particularly preferred combinations are the same as Preferred integers, groups or combinations and

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Preferred compounds having Formula Vb are:

MeOOC-CO-(R)Phe-Phe-Pab(Z) Boc-(R)Phe-Phe-Pab(Z)  $Bnooc-CH_2-(R)Pro-Phe-Pab(Z)$  $BnOOC-CH_2-(R)$  Phe-Phe-Pab(2) Boc-(R)Pro-Phe-Pab(Z)

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use of a compound of the formula; In a further embodiment the invention relates to novel 25

kininogenases inhibitors. It can be used as such or synthesis of peptidic thrombin inhibitors or serine protease inhibitor, and in particular in as a starting material in synthesis of a peptidic

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Pharm. vol 23, p. 2247-2256. has been previously disclosed in inter alia Biochem. aminomethylbenzene " or "H-Pab" herein. The compound compounds. This compound is named "1-amidino-4carried out by methods known in the art for amidino carbonyl. Protection of the amidino derivatives is the nitrogens with a protective group such as benzyloxy having the amidino group either mono- or diprotected at

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6 The structural fragment of the formula

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20 a thrombin inhibitor or kininogenases inhibitor renders a serine protease inhibitor, and in particular compound, especially a peptic compound. The fragment structural element in a pharmaceutically active has however not been previously disclosed as a

25 use of a compound of the formula: In a further embodiment the invention relates to novel

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known in the art for amidino compounds. This compound protective group such as benzyloxy carbonyl. Protection of the amidino derivatives is carried out by methods either mono- or diprotected at the nitrogens with a inhibitor. The compound may have the amidino group as a starting material in synthesis of a thrombin

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is named "1-amidino-4-aminomethyl cyclohexane" or The compound has been previously disclosed in DE "H-Pac" herein.

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The structural fragment of the formula

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structural element in a thrombin inhibitor valuable. has however not been previously disclosed as a

In a further embodiment the invention relates to a novel compound of the formula:

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derivatives is carried out by methods known in the art and the use of said compound as a starting material in synthesis of a serine protease inhibitor, especially a such as beneyloxy carbonyl. Protection of the amidino diprotected at the nitrogens with a protective group compound may have the amidino group either mono- or aminoethyl-1-amidino piperidine" or "H-Rig" herein. thrombin inhibitor or kininogenase inhibitor. The for amidno compounds. This compound is named "4-30 25

The structural fragment of the formula

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renders a serine protease inhibitor, and in particular compound, especially a peptic compound. The fragment structural element in a pharmaceutically active a thrombin inhibitor or kininogenases inhibitor has however not been previously disclosed as a

In a further embodiment the invention relates to a novel compound of the formula:

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varuable.

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derivatives is carried out by methods known in the art and the use of said compound as a starting material in such as benzyloxy carbonyl. Protection of the amidino synthesis of a serine protease inhibitor especially a diprotected at the nitrogens with a protective group for amidino compounds. This compound is named "1,3compound may have the amidino group either mono- or diaza-2-imino-4-aminoethyl cyclohexane" or "H-Itp" thrombin inhibitor or kininogenase inhibitor. The herein.

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The structural fragment of the formula

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has however not been previously disclosed as a strucural element in a pharmaceutically active

a thrombin inhibitor or kiniogenases inhibitor renders a serine protease inhibitor, and in particular compound, especially a peptic compound. The fragment

compounds of the formula: In a further embodiment the invention relates to novel

where n is 1 or 2 s is 0 ro 1,

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for amidino compounds. These compounds are named: derivatives is carried out by methods known in the art such as benzyloxy carbonyl. Protection of the amidino diprotected at the nitrogens with a protective group compound may have the amidino group either mono- or thrombin inhibitors or kininogenases inhibitors. The synthesis of serine protease inhibitors, especially and the use of said compounds as a starting material in

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is 1 and s is 1 1-amidino-3-aminomethyl pyrrolidine or "H-Nig" when n 25

3-aminomethyl-1-amidino azetidine or "H-Mig" when n is 2 and s is 1 1-amidino-3-aminoethyl pyrrolidine or "H-Hig" when n is

and s is 0 3-aminoethyl-1-amidino azetidine or "H-Dig" when n is 2 1 and s is 0

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The structural fragment of the formula

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15 a thrombin inhibitor or kininogenases inhibitor renders a serine protease inhibitor, and in particular compound, especially a peptic compound. The fragment structural element in a pharmaceutically active has however not been previously disclosed as a

20 compounds having the amidino group mono- or digroup, examples of such compounds are protected at the nitrogens with a benzyloxy carbonyl A further embodiment of the invention are the novel

25 (H-Pab(Z)), 4-aminomethyl-1-(N-benzyloxycarbonylamidino) benzene

benzene  $(H-Pab(Z)_2)$ , 4-aminomethyl-1-(N,N'-di(benzyloxycarbonyl)amidino)

cyclohexane (H-Pac(Z)), 4-aminomethyl-1-(N-benzyloxycarbonylamidino)

cyclohexane  $(H-Pac(Z)_2)$ , 4-aminomethyl-1-(N,N'-di(benzyloxycarbonyl)amidino)

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4-aminoethyl-1-(N-benzyloxy-carbonylamidino piperidine

piperidine (H-Rig(Z)2), (3RS)-1-(N-benzyloxycarbonylamidino)-3-aminomethyl 4-aminoethyl-1-N,N'-di(benzyloxycarbonyl)amidino

pyrrolidine (H-Nig(2)),

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(3RS)-1-(N-benzyloxycarbonylamidino)-3-aminoethyl (3RS)-1-(N,N'-di(benzyloxycarbonyl)amidino)-3aminomethyl pyrrolidine (H-Nig( $\mathbb{Z})_2$ ), pyrrolidine (H-Hig(Z)),

3-aminomethyl-1-(N-benzyloxycarbonylamidino) azetidine (3RS)-1-(N,N'-di(benzyloxycarbonyl)amidino)-3aminoethyl pyrrolidine (H-Hig( $^{\rm Z}$ ),

3-aminoethy1-1-(N-benzyloxycarbonylamidino) azetidine 3-aminomethyl-1-(N,N'-di(benzyloxycarbonyl)amidino) azetidine  $(H-Mig(Z)_2)$ , (H-Mig(Z)),

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3-aminoethyl-1-(N,N'-di(benzyloxycarbonyl)amidino) azetidine  $(H-Dig(Z)_2)$ , (H-Dig(Z)),

Said compounds are used as starting materials in the preparation of the claimed peptide derivatives of formulas I, Ia, Ib, V, Va and Vb.

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### Medical and pharmaceutical use 20

conditions where inhibition of thrombin is required and for the treatment, in a human or animal organism, of of physiologically disorders especially inflammatory The invention also provides compositions and methods diseases.

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treatment and/or prophylaxis, include venous thrombosis including man in treatment or prophylaxis of thrombosis Alzheimers disease and pancreatitis. Disease states in there is an undesirable excess of the thrombin without The thrombin inhibiting compounds of the invention are and hypercoagulability in blood and tissues. They are furthermore expected to be useful in situations where which these compounds have a potential utility, in signs of hypercoagulability, for example as in expected to be useful in particular in animals

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in haemodialysis.

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arterial fibrillation or from the left ventricule after and pulmonary embolism, arterial thrombosis, such as in based stroke and peripheral arterial thrombosis and myocardial infarction, unstable angina, thrombosissystemic embolism usually from the atrium during

atherosclerotic diseases such as coronary arterial

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compounds have expected utility in prophylaxis of

transmural myocardial infarction. Further, these

general. Further, these compounds have expected utility intravascular coagulation caused by bacteria, multiple combined with any antithrombotic agent with a different arterial disease. Further, these compounds are expected anticoagulant treatment when blood is in contact with cardiovascular surgery using or heart-lung machine or anticoagulant treatment when blood is in contact with thrombosis after microsurgery and vascular surgery in (PTCA) and coronary bypass operations. Further, these trauma, intoxication or any other mechanism. Further, foreign surfaces in the body such as vascular grafts, vasculars stemts, vascular catheters, mechanical and thrombolysis, percutaneous trans-luminal angioplasty compounds have expected utility in prevention of reexpected to be useful together with thrombolytics in mechanism of action, such as the antiplatelet agent biological prosthetic or any other medical device. Further, these compounds have expected utility in infarction. Further, these compounds have expected acetylsalicylic acid. Further, these compounds are disease, cerebral arterial disease and peripheral medical devices outside the body such as during to have synergistic antithrombotic effects when chrombotic diseases, in particular myocardial utility in prophylaxis for re-occlusion after these compounds are expected to be useful in in treatment and prophylaxis of disseminated 30 25 20 15

other blood products in vitro. A further expected utility of the anticoagulant anticoagulants for preservation of blood, plasma and and mechanical devises used in patients in vivo, and as compounds of the invention are in rinsing of catheters

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combination with other therapeutic agents. acceptable carrier or diluent can be used solely or in inhibiting compounds with or without a physiologically and arthritis. An effective amount of kininogenase pancreatitis, uticaria, inflammatory bowel diseases, inflammatory diseases such as asthma, rhinitis, animals including man in treatment or prophylaxis of invention are expected to be useful in particular in The antiinflammatory inhibiting compounds of the

processes in airway mucosa or gut mucosa. present compounds can for example be studied by their assessed with chromogenic substrates according to known inhibition of allergen-induced exudative inflammatory procedures. The anti-inflammatory actions of the The compounds inhibit the activity of kallikreins

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### Pharmaceutical preparations

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pharmaceutically acceptable dosage form. Depending upon tartrate, trifluoroacetate and the like in a lactate, acetate, citrate, bensoate, succinate, hydrobromide, sulphate, hydrosulphate, nitrate, or a pharmaceutical acceptable non-toxic organic or route, in the form of pharmaceutical preparations comprising the active ingredient either as a free base tracheally, bronchially, parenterally or via inhalation administered orally, rectally, dermally, nasally, inorganic acid addition salt, e.g. the hydrochloride, The compounds of the invention will normally be

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administration, the compositions may be administered at the disorder and patient to be treated and the route of

varying doses.

suitable for oral administration. between 0.2 and 75 % by weight for preparations preparations intended for parenteral administration and 0.1 and 99 % by weight of the preparation, more specifically between 0.1 and 50 % by weight for preparation prepared by per se known techniques. Usually the active substance will constitute between The dosage form may be a solid, semisolid or liquid

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100 mg/kg body weight at peroral administration and administration. 0.001-50 mg/kg body weight at parenteral in therapeutical treatment of humans are about 0.001-Suitable daily doses of the compounds of the invention

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#### Preparation

25 methods to a compound second aminoacid is added afterwards using standard aminoacid, when a N-terminally amino acid is used a coupling of an N-terminally protected dipeptide or Formula I and V may be prepared by processes comprise preparation of the compounds. The compounds of A further objective of the invention is the mode of

H<sub>2</sub>N----(CH<sub>2</sub>)<sub>n</sub>-----X

35 in formula V as such or having the guanidino or amidino where B is as defined in formula I and D is as defined wherein n is an integer 0, 1, 2, 3 or 4, X is B or B-D

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butyloxy carbonyl- or p-toluenesulphonyl- group or X is salt, and in those cases where the reaction results in re-crystallisation techniques, and if desired a single a group transferable into B followed by removal of the protectary group(s) or deprotection of the N-terminal nitrogen and if desired deprotection by known methods protecting group such as a benzyloxy carbonyl-, tertand if desired forming a physiologically acceptable nitrogen followed by alkylation of the N-terminal nitrogens either mono or diprotected with an amin a mixture of stereoisomers, these are optionally separated by standard chromatographic or

In more detail the compounds of Formula I or V may be prepared by either of the following methods:

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stereoisomer is isolated.

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#### Method Ia

selected from  ${\mathtt A}^1$  and  ${\mathtt A}^2$  in Formulas I or V and prepared Coupling of an N-terminally protected dipeptide, by standard peptide coupling, with a compound 20

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using standard peptide coupling, shown in the formula

$$-A^1 - A^2 - \omega$$

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$$\begin{pmatrix} H_2N-(CH_2)^n & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & \\ & & & \\ & & \\ & & & \\ & &$$

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when  $\mathbb{Q}^1$  is -NH-W<sup>2</sup> (W<sup>2</sup> in this case must be orthogonal to carbonyl or benzyloxy carbonyl, or  $\mathbb{Q}^1$  is -CN, -CO-NH $_2$  or subsequently transferred into a guanidino group (giving C(NW<sup>2</sup>)-NH-W<sup>2</sup>, -C(NH)-NH-W<sup>2</sup>, -NH-C(NH)-NH<sub>2</sub>, -NH-C(NH)-NH- $W^2$  ,  $-N\left(W^2\right)-C\left(NH\right)-NH-W^2$  or  $-NH-C\left(NW^2\right)-NH-W^2$  , where  $W^2$  is wherein n is as defined in Formula I  $\mathtt{W}^1$  is an N-teminal  $\boldsymbol{Q}^1 = -NH - C\left(NH\right) - NH_2\right)$  , after deprotection of the  $W^2 - g \mathbf{roup}$ methods known in the art or  $\mathbb{Q}^1$  is  $\mathrm{NH}_2$  or  $\mathrm{NH}\text{-W}^2$ , where amino protecting group such as tert-butyloxy carbonyl -cs-NH $_{
m 2}$ , where the group is subsequently transferred into a amidino group (e.g giving  $\mathbf{Q}^1 = - \mathbf{C}(\mathrm{NH}) - \mathrm{NH}_2)$  by  $W^2$  is as defined above, where the amino group is and benzyloxy carbonyl and and  $\mathbb{Q}^1$  is  $-\text{C(NH)}-\text{NH}_2,$ an amine protecting group such as tert-butyloxy

 $C(NH)-NH-W^2$  or  $-NH-C(NW^2)-NH-W^2$  (W^2 in this case must be Removal of the protecting group(s) (when  $\mathbb{Q}^{1_m} - \mathbb{C}(\mathrm{NH}) - \mathrm{NH}_2$ , The final compounds can be made in any of the following -C(NW<sup>2</sup>)-NH-W<sup>2</sup>, -C(NH)-NH-W<sup>2</sup>' -NH-C(NH)-NH<sub>2</sub>' -NH-C(NH)selective deprotection of the  $W^{1-}$  group (e.g when  $\mathbb{Q}^{1=}$ terminal nitrogen by methods known in the art and if  $\mathrm{NH-W}^2$ ,  $-\mathrm{N}(\mathrm{W}^2)-\mathrm{C}(\mathrm{NH})-\mathrm{NH-W}^2$  or  $-\mathrm{NH-C}(\mathrm{NW}^2)-\mathrm{NH-W}^2)$ , or a ways, depending on the nature of the  $\mathbf{Q}^1-$  group used:  $C(NW^2)$  -NH- $W^2$ , -C(NH) -NH- $W^2$ , -NH-C(NH) -NH- $W^2$ , -N( $W^2$ ) orthogonal to  $W^1$ ) followed by alkylation of the Ndesired deprotection by known methods.

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 $\ensuremath{\text{W}}^1$  ), by methods known in the art.

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#### Method Ib

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acid, selected from  $\mathbb{A}^2$  in Formulas I or V and prepared by standard methods, with a compound of formula Coupling of an N-terminally protected amino 30

using standard peptide coupling, shown in the formula

 $w^1 - A^2 - OH$ 

$$H_2N - (CH_2) \frac{1}{n}$$

 $W^1$ — $A^2$ —HN— $(CH_2)$ n

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wherein n,  $w^1$ , and  $Q^1$  are as defined above followed by deprotection of the  $w^1$ -group and coupling with the N-terminal amino acid, in a protected form, leading to the protected peptide described in Method Ia. The synthesis to the final peptides is then continued according to Method Ia.

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#### Method IIa

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by standard peptide coupling, with a compound

Coupling of an N-terminally protected dipeptide, selected from  $\mathbb{A}^1$  and  $\mathbb{A}^2$  in Formulas I or V and prepared

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using standard peptide coupling, shown in the formula

$$\begin{array}{c}
W' - A' - A^2 - QH \\
H_2N - (CH_2)_n - Q^1
\end{array}$$

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30 25 20 orthogonal to  $W^1$  ), by methods known in the art. the  $W^2$ -group when  $Q^1$  is  $-NH-W^2$  ( $W^2$  in this case must be group (giving  $Q^{1_{\pm}}$  -NH-C(NH)-NH $_{2}$ ), after deprotection of group is subsequently transferred into a guanidino  $^{
m NH-W^2}$ , where  $^{
m W^2}$  is as defined above, where the amino  $C(NH)-NH_2)$  by methods known in the art or  $Q^1$  is  $NH_2$  or transferred into a amidino group (e.g giving  $Q^{1}=$  - $^{
m CN, -CO-NH_2}$  or  $^{m CS-NH_2, \ where the group is subsequently}$ butyloxy carbonyl or benzyloxy carbonyl, or  $Q^1$  is where  $W^2$  is an amine protecting group such as tert- $C(NH)-NH-W^2$ ,  $-N(W^2)-C(NH)-NH-W^2$  or  $-NH-C(NW^2)-NH-W^2$ ,  $^{\rm NH}_2$ ,  $^{\rm -C}(^{\rm NW}^2)$   $^{\rm -NH-W}^2$ ,  $^{\rm -C}(^{\rm NH})$   $^{\rm -NH-W}^2$ ,  $^{\rm -NH-C}(^{\rm NH})$   $^{\rm -NH}_2$ ,  $^{\rm -NH-W}$ carbonyl and benzyloxy carbonyl and and  $Q^1$  is -C(NH)teminal amino protecting group such as tertbutyloxy wherein n is as defined in Formula I,  $w^1$  is an N-

The final compounds can be made in any of the following ways, depending on the nature of the Q<sup>1</sup>- group used: Removal of the protecting group(s) (when Q<sup>1</sup>- -C(NH)-NH<sub>2</sub>, -C(NH)-NH-W<sup>2</sup>, -NH-C(NH)-NH<sub>2</sub>, -NH-C(NH)-NH<sub>2</sub>, -NH-C(NH)-NH-W<sup>2</sup>, -N(W<sup>2</sup>)-C(NH)-NH-W<sup>2</sup> or -NH-C(NW<sup>2</sup>)-NH-W<sup>2</sup>), or a selective deprotection of the W<sup>1</sup>- group (e.g when Q<sup>1</sup>= -

C(NW<sup>2</sup>)-NH-W<sup>2</sup>, -C(NH)-NH-W<sup>2</sup>, -NH-C(NH)-NH-W<sup>2</sup>, -N(W<sup>2</sup>)-C(NH)-NH-W<sup>2</sup> or -NH-C(NW<sup>2</sup>)-NH-W<sup>2</sup> (W<sup>2</sup> in this case must be orthogonal to W<sup>1</sup>) followed by alkylation of the N-terminal nitrogen by methods known in the art and if desired deprotection by known methods.

#### Method IIb

Coupling of an N-terminally protected amino acid, selected from  $A^2$  in Formulas I or V and prepared by standard methods, with a compound of formula

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using standard peptide coupling, shown in the formula

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$$w' - A^{2} - cH$$

$$\downarrow^{H_{2}N - (CH_{2})} - CH$$

$$w' - A^{2} - HN - (CH_{2}) - CH$$

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wherein n, W<sup>1</sup> and Q<sup>1</sup> are as defined above followed by deprotection of the W<sup>1</sup>-group and coupling with the N-terminal amino acid, in a protected form, leading to the protected peptide described in Method IIa. The synthesis to the final peptides is then continued according to Method IIa.

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#### Method IIIa

Coupling of an N-terminally protected dipeptide, selected from  ${\bf A}^1$  and  ${\bf A}^2$  in Formulas I or V and prepared by standard peptide coupling, with a compound

$$H_2N - (CH_2) \frac{(CH_2)^2 - X^4}{N}$$

using standard peptide coupling, shown in the formula

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wherein n is as defined in Formula I and r is 0.1 when  $x^1$ ,  $x^2$  and  $x^4$  are  $CH_2$  or r is 0 when  $x^2$  and  $x^4$  are  $CH_2$  and  $x^1$  is abscent,  $w^1$  is an N-teminal amino protecting group such as tert-butyloxy carbonyl and benzyloxy carbonyl and and  $Q^2$  is  $-C(NH)-NH_2$ ,  $-C(NW^2)-NH-W^2$ , or  $-C(NH)-NH-W^2$ , where  $W^2$  is an amine protecting group such as tert-butyloxy carbonyl or benzyloxy carbonyl, or  $Q^2$  is equal to  $W^2$  where the amino group, after deprotection of the  $W^2$  group  $(W^2$  in this case must be orthogonal to  $W^1$ , is subsequently transferred into a guanidino group using a unprotected, N-protected or N,N'-diprotected guanidation reagent by methods known

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in the art (giving Q2= -C(NH)-NH2, -C(NW2)-NH-W2 or -C(NH)-NH-W2).

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The final compounds can be made in any of the following ways, depending on the nature of the  $Q^2$ - group used: Removal of the protecting group(s) (when  $Q^2$ -  $-C(NH)-NH_2$ ,  $-C(NW^2)-NH-W^2$  or  $-C(NH)-NH-W^2$ ), or a selective deprotection of the  $W^1$ - group (e.g when  $Q^2$ -  $-C(NW^2)-NH-W^2$ ,  $-C(NH)-NH-W^2$ ,  $W^2$  in this case must be orthogonal to  $W^1$ ) followed by alkylation of the N-terminal nitrogen by methods known in the art and if desired deprotection known methods.

#### Method IIIb

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Coupling of an N-terminally protected amino acid, selected from  $\mathbb{A}^2$  in Formulas I or V and prepared by standard methods, with a compound of formula

$$H_2N \longrightarrow (CH_2) \bigcap_{n \longrightarrow X^2} (CH_2) \bigcap_{n \longrightarrow X^2} X^1 \longrightarrow X^2$$

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using standard peptide coupling, shown in the formula

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wherein n, r,  $x^1$ ,  $x^2$  and  $x^4$ ,  $y^1$ , and  $Q^2$  are as defined

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above followed by deprotection of the w<sup>1</sup>-group and coupling with the N-terminal amino acid, in a protected form, leading to the protected peptide described in Method IIIa. The synthesis to the final peptides is then continued according to Method IIIa.

#### Method IVa

Coupling of an N-terminally protected dipeptide, selected from  ${\bf A}^1$  and  ${\bf A}^2$  in Formulas I or V and prepared by standard peptide coupling, with a compound

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using standard peptide coupling, shown in the formula

wherein n is as defined in Formula I, W<sup>1</sup> is an N-terminal amino protecting group such as tert-butyloxy carbonyl or benzyloxy carbonyl and W<sup>3</sup> is H or an amino protecting group such as aryl sulfonyl, benzyloxy carbonyl or tert-butyloxy carbonyl. The final compounds

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can be made in any of the following ways: Removal of the protecting group(s), or a selective deprotection of the W<sup>1</sup>-group (W<sup>1</sup> must be orthogonal to W<sup>3</sup>) followed by alkylation of the N-terminal nitrogen and if desired deprotection.

Method IVb

Coupling of an N-terminally protected amino acid, selected from A<sup>2</sup> in Formulas I or V and prepared by standard methods, with a compound of formula

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using standard peptide coupling, shown in the formula

W.

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$$w' - A^2 - v_H - (cH_2) \frac{NH}{N - w^3}$$

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wherein n,  $W^1$ , and  $W^3$  are as defined above followed by deprotection of the  $W^1$ -group ( $W^1$  must be orthogonal to  $W^2$ ) and coupoing with the N-terminal amino acid, in a protected form, leading to the protected peptide described in Method IVa. The synthesis to the final peptides is then continued according to Method IVa.

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## DETAILED DESCRIPTION OF THE INVENTION

The following description is illustrative of aspects of the invention.

EXPERIMENTAL PART

General experimental Procedures.

10 Mass spectra were recorded on a Finnigan MAT TSQ 700 triple guadropole mass spectrometer equipped with an electrospray interface.

The <sup>1</sup>H NMR and <sup>13</sup>C NMR measurements were performed on BRUKER AC-P 300 and BRUKER AM 500 spectrometers, the former operating at a <sup>1</sup>H frequency of 500.14 MHz and a <sup>13</sup>C frequency of 125.76 MHz and the latter at <sup>1</sup>H and <sup>13</sup>C frequency of 300.13 MHz and 75.46 MHz respectively.

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The samples were about 10-50 mg dissolved in 0.6 ml of either of the following solvents; CDCl<sub>3</sub> (isotopic purity > 99.8%), CD<sub>3</sub>OD (isotopic purity > 99.9%), D<sub>3</sub>OD (isotopic purity > 99.9%) or DMSO-d<sub>6</sub> (isotopic purity < 99.8%). All solvents where purchased from Dr. Glaser AG, Basel.

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The  $^1\text{H}$  and  $^{13}\text{C}$  chemical shift values in CDCl<sub>3</sub> and CD<sub>3</sub>OD are relative to tetramethylsilane as an external standard. The  $^1\text{H}$  chemical shifts in D<sub>2</sub>O are relative to the sodium salt of 3-(trimethylsilyl)-d<sub>4</sub>-propanoic acid and the  $^{13}\text{C}$  chemical shifts in D<sub>2</sub>O are referenced relative to 1,4-dioxane (67.3 ppm), both as external relative to 1,4-dioxane with an external standard may in

some cases cause minor shift differences compared to an internal standard, however, the difference in  $^1{\rm H}$  chemical shift is less than 0.02 ppm and in  $^{13}{\rm C}$  less than 0.1 ppm.

existence of two contributing conformers with respect proline or a "proline like" residue frequently exhibits two sets of resonances. This corresponds to the The  $^1H$  NMR spectrum of peptide sequences containing a

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than the number of protons expected from the chemical protons reported for some of the intermediates are less NMR-documentation. This implies that the number of formula. signals are clearly resolved they are reported in the NH-signals in CDCl3, only in the cases where the NMR documentation. The same criterium is valid for the Only in the cases where the signals of the minor rotamer are clearly resolved they are reported in the  $^{
m l}$ H chemical shifts of the major rotamer is reported. cis-trans equilibrium with one conformer as the preponderant conformer (>90%). In those cases only the Aze-, (R)Cha-Pro- and (R)Cha-Pic- often give rise to a cis and trans. In our compounds the sequences (R)Chathe N-part of the amide bond. The conformers are named to the rotation around the amide bond, where proline is

followed by spraying with a solution prepared by mixing Visualisation was by a combination of UV-light, Merck Silicagel 60 $F_{254}$  coated glass or aluminium plates Thin-Layer Chromatography was carried out on commercial

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gel 60 (40-63 mm, 230-400 mesh) under pressure of air. Flash chromatography was carried out on Merck Silica % in EtOH(95%)) and heating. ml of concentrated acetic acid and 10.2 ml of p-methoxy benzaldehyde or phosphomolybdic acid reagent (5~10 w.t 372 ml of EtOH(95%), 13.8 ml of concentrated  $\mathrm{H}_2\mathrm{SO}_4$ , 4.2

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Reversed phase high-performance liquid chromatography

phase Kromasil 100,C8 columns (Eka-Nobel) having a Waters M-590 instrument equipped with three reverse (in the Examples referred to as RPLC) was performed on

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mm) chromatography detecting at 226 nm. different dimensions for analytical (4.6 mm x 250 mm), semipreparative (1  $^{\circ}$  x 250 mm) and preparative ( 2  $^{\circ}$  x 500

Lyovac GT 2, apparatus. Freeze-drying was done on a Leybold-Heraeus, model

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### Preparation of starting materials

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Boc-(R) Pg1-OH

Prepared in the same way as described for Boc-(R)Cha-OH (vide infra) from H-(R)Pgl-OH.

#### Boc-(R)Cha-OH

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25 20 solvent afforded 30.9 g (90.5 %) of the title compound water, brine and dried  $(Na_2SO_4)$ . Evaporation of the of EtOAc. The combined organic phase was washed with acidified with 2 M KHSO $_4$  and extracted with 3 x 150 ml aqueous phase was washed twice with EtOAc, then at room temperature. The THF was evaporated and an To a solution of H-(R)Cha-OH, 21.55 g (125.8 mmol), in as a white solid. additional 150 ml of water was added. The alkaline mmol) of (Boc)20 and the mixture was stirred for 4.5 h 130 ml 1 M NaOH and 65 ml THF was added 30 g (137.5

#### 30 Boc-(R) Hop-OH

(R) Cha-OH starting from H-(R) Hop-OH. Prepared by the same procedure as described for Boc-

ដូ 1H), 7.15-7.33 (m, 5H). 2.22 (m, 1H), 2.75 (bt, 2H), 4.36 (bs, 1H), 5.05 (bs, <sup>1</sup>H~NMR (300 MHz, CDCl<sub>3</sub>): & 1.45 (s, 9H), 2.00 (m, 1H),

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# 4-(tert-butyloxycarbonylaminomethyl) pyridine

on standing. The crude product was used without further filtered through silica gel. Evaporation of the solvent gave the title compound as a red oil which crystallized pyridine in 100 ml THF was added 24 g (110 mmol)  $\mathrm{Boc}_2\mathrm{O}$ stirred for 4 h (a precipitate was formed during the reaction and the slurry became red).The solvent was solution was allowed to reach room temperature and removed and the residue was dissolved in EtoAc and dissolved in 70 ml THF at 10°C for 20 minutes. The To a solution of 10.81 g (100 mmol) 4-aminomethyl purification.

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1H-NHR (300 MHz, CDCl<sub>3</sub>): 6 1.45 (s, 9H), 4.32 (d, 2H), 5.05 (bs, 1H (NH)), 7.2 (d, 2H), 8.55 (d, 2H). 15

4-aminomethy1-1-(N-benzyloxycarbonylamidino)-benzene (H-Pab(Z))

(i) 4-cyanobenzyl azide

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phase was extracted with an additional 2x50 ml toluene. water and brine and finally dried (MgSO $_{f 4}$ ) and filtered. The combined organic extracts were washed with 2x50 ml potentially explosive azide compounds it is adviceable A solution of 20.23 g (0.31 mol) sodium azide in 50 ml addition of the water) and 500 ml water. The aqueous water was added to 49.15 g (251 mmol) 4-cyanobenzyl exothermic reaction took place and after 1.5 h the to add the toluene to the rection mixture before bromide in 200 ml DKF at ambient temperature. An toluene(caution: In order to avoid separation of The solution was used as such in the next step. reaction mixture was diluted with 200 ml

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1H-NMR (300 MHz, CDCl3); 6 4.4 (s, 2H), 7.4 (d, 2H), 7.7

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(d, 2H).

(ii) 4-amidino benzyl azide

this point the yield of 4-amidino benzyl azide was 22.5 anhydrous ether gave white crystals which were isolated crystals. The crystals were isolated by filtration. At and 200 ml 3.75 M NaOH solution was added whereupon 4-Storage at 8°C for 24 h and evaporation of most of the Hydrogen chloride was bubbled into a mixture of 250 ml (approximatly 200 ml) above at - 5°C until saturation. ammonia. After 48 h most of the solvent was removed by filtration and dissolved in 1.8 l of alcoholic solvent followed by precipitation by addition of amidino benzyl azide precipitated as colourless absolute ethanol and the solution from step (i)g (total 51%). 12 9

Ethylimidatobenzyl azide hydrochloride:

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4.65 (q, 2H), 4.8 (br s, 2H), 7.6 (d, 2H), 8.1 (d, 2H) H-NWR (500 MHz, CD<sub>3</sub>OD); & 1.6 (t, 3H), 4.5 (s, 2H),

4-amidino benzyl azide:

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<sup>1</sup>H-NYR (500 MHz, CDCl<sub>3</sub>); 6 4.3 (s, 2H), 5.7 (br s, 3H), 7.3 (d, 2H), 7.6 (d, 2H).

13C-NMR (125 MHz, CDCl<sub>3</sub>): amidine carbon: 6 165.5.

(iii) 4-(benzyloxycarbonylamidino) benzyl azide

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methylene chloride and the resulting solution was dried was added. 25 ml Benzyl chloroformate was slowly added to the stirred solution while the reaction mixture was  $(K_2CO_3)$ , filtered and 27 ml (194 mmol) triethyl amine The crystals from (ii) above were dissolved in 500 ml

with 2M HCl. The organic phase was dried (MgSO $_{\rm 4}$ ) and chloride/hexane. isolated as colorless crystals from ether/methylene (benzyloxycarbonylamidino) benzyl azide was finally the solvent was removed in vacuo. 4was added and the aqueous phase was adjusted to pH  $7\,$ continued for another 30 minutes. Subsequently, water ml benzyl chloroformate was added and stirring was cooled in an ice bath. After 30 minutes an additional 2

6.3-7.0 (br s, 1H), 7.3-7.4 (m, 5H), 7.5 (d, 2H), 7.9 1H-NMR (500 MHz, CDCl<sub>3</sub>); & 4.4 (s, 2H), 5.3 (s, 2H), (d, 2H), 9.3-9.6 (br s, 1H).

15  $^{13}\text{C-NMR}$  (125 MHz, CDCl<sub>3</sub>): amidine carbon: & 167.5.

benzene (H-Pab(Z)) (iv) 4-aminomethyl-1-(N-benzyloxycarbonylamidino)-

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yellow oil which solidified on standing. yield starting from cyanobenzyl bromide is 28%) of a methylene chloride followed by drying  $(K_2CO_3)$  and with 3.75M sodium hydroxide solution. Extraction with removal of the solvent in vacuo gave 20 g (The total chloride and ether and was subsequently made alcaline HCl. The aqueous phase was washed with methylene dissolved in methylene chloride and extracted with 2M before removal of the solvent in vacuo. The residue was was added and the solution was allowed to stand for 4 h  $\,$ 16 h an additional 6.6 g (25 mmol) triphenylphosphine azide from (iii) above dissolved in 160 ml THF. After temperature to the 4-(benzyloxycarbonylamidino) benzyl 26.3 g (100 mmol) triphenylphosphine was added at room

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1H-NMR (500 MHz, CDCl3); & 1.2-2.2 (br s, 2H), 3.8 (s, 2H), 5.2 (s, 2H), 7.2-7.35 (m, 5H), 7.4 (d, 2H), 7.8

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(d, 2H), 9.1-9.6 (br s, 1H).

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δ 164.6 and 168.17.  $^{13}\mathrm{C\text{-}NMR}$  (125 MHz, CDCl $_3$ ): amidine and carbonyl carbons:

H-Pig(2)2

(i) 4-(tert-butyloxycarbonyl-aminomethyl) piperidine

30 25 20 15 10 and evaporated to give 5.2 g of the title compound as a white powder. + 1 imes 75 ml EtOAc. The combined organic phase was dried made alkaline with 2 M NaOH and extracted with 1 imes 200  $m1 + 1 \times 25 m1 1 M KHSO_4$ . The combined acidic phase was ml EtOAc and the organic phase was washed with 1 imes 50 of a white oil. The brown powder was dissolved in 100 brown powder. Evaporation of the mother liqour gave 7  ${\sf g}$ a precipitate which was filtered off to give 7.7 g of a 50 ml of diethyl ether. Addition of 200 ml pentane gave gave a 17.2 g of a brownish oil which was dissolved in water and dried (MgSO $_4$ ). Evaporation of the solvent  $\mathrm{CH_2Cl_2}.$  The combined organic phase was washed with 25 ml water phase was extracted with 1  $\times$  200 + 1  $\times$  100 ml the mixture was made alkaline with 5 M NaOH and the vaccuo. After addition of 50 ml water to the residue filtered off and most of the acetic acid was removed in hydrogenated for 4 days at 0.34 MPa. The catalyst was acid, 2 g of 5  $Rh/Al_2O_3$  was added and the mixture was vaccuo and the residue was dissolved in 100 ml acetic catalyst was filtered off and the solvent removed in the hydrogenation was incomplete. Therefore, the hydrogenated at 0.34 MPa over night.  $^1\mathrm{H-NMR}$  showed that added 2 g of 5 %  $\mathrm{Rh/Al_2O_3}$  and the mixture was butyloxycarbonylaminomethyl pyridine in 125 ml MeOH was To a solution of 17.7 g 4-tert-

Treatment of the white oil obtained from the mother

ligour above in the same way afforded an additional 3.4 g of the product. Total yield 40 %.

major rotamer: 6 1.11 (dg, 2H), 1.44 (s, 9H), 1.49-1.60  $^{1}\mathrm{H-NMR}$  (500 MHz, CDCl3, mixture of two rotamers, 3:1): (m, 1H), 1.63-1.70 (m, 2H), 2.58 (dt, 2H), 2.93-3.03 (m, 2H), 3.07 (m, 2H), 4.75 (bs, 1H (NH)).

Resolved signals arising from the minor rotamer appear at 6 1.21 (dg) and 1.91 (dt).

### (ii) Boc-Pig(Z)<sub>2</sub>

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butyloxycarbonyl-aminomethyl) piperidine in 60 ml  ${
m CH_3CN}$ pethroleum ether/EtOAc (1/1) as eluent afforded 2.43 g carbonyl)methylisothiourea and the mixture was stirred residue was dissolved in EtOAc. The organic phase was washed with 2 x 20 ml 1 M KHSO4, 1 x 20 ml water, 1 x at 60°C for 22 h. The solvent was evaporated and the was added 3.34 g (9.33 mmol) of N,N'-(dibenzyloxy-20 ml brine and dried(MgSO $_4$ ). Evaporation of the solvent followed by flash chromatography using To a solution of 2 g (9.33 mmol) 4-(tert-(50%) of the desired product. 20

H-NMR (500 MHz, CDCl<sub>3</sub>): Some signals, especially in the 2.66-3.05 (m, 4H), 3.7-4.5 (bs, 2H), 4.65 (bt, 1H(NH)), 6 1.19-1.31 (m, 2H), 1.43 (s, 9H), 1.63-1.80 (m, 3H), piperidine ring, are selectively broadend due to an intramplecular exchange process. This is especially piperidine ring, which exhibit a broad peak ranging 5.13 (8, 4H), 7.2-7.4 (m, 10H), 10.5 (bs, 1H(NH)). pronounced for the 2- and 6-CH $_2$  groups of the from 3.7 to 4.5 ppm.

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(111) H-Pig(Z)<sub>2</sub>

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 $\mathtt{CH_2Cl_2}$ . The organic phase was washed with 5 ml 2 M NaOH, Evaporation of the solvent afforded 100 mg (76 %) of A solution of 163 mg (0.31 mmol) Boc-Pig( $\mathbf{2}$ ) in 5 ml EtOAc saturated with HCl(g) was stirred at ambient temperatur for 3 h and 20 minutes. The solvent was evaporated and the residue was dissolved in 30 ml 1 x 5 ml water, 1 x 5 ml brine and dried(MgSO $_{4}$ ).

<sup>1</sup>H-NWR (500 MHz, CDCl<sub>3</sub>): Some signals, especially in the piperidine ring, which exhibit a broad peak ranging intramolecular exchange process. This is especially piperidin ring, are selectively broadend due to an pronounced for the 2- and 6- $\mathrm{CH}_2$  groups of the from 3.7 to 4.5 ppm. 2 52

the title compound.

2H), 2.57 (d, 2H), 2.86-3.03 (m, 2H), 3.7-4.5 (bs, 2H), 6 1.18-1.37 (m, 2H), 1.46-1.63 (m, 1H), 1.68-1.83 (m, 5.13 (s, 4H), 7.2-7.4 (m, 10H).

4-aminomethyl-1-(N-benzylowy carbonylamidino)cyclohexane (H-Pac(Z) x 2HCl). 20

(i) N-[N-4-(benzyloxycarbonyl)amidino benzyl] tertbutyl carbamate

mmol) triethyl amine in 25 ml methylene chloride. After (benzyloxycarbonyl)amidino benzyl amine and 1 ml (7.1 mixture was washed with 5% acetic acid and 10% sodium 1.466 g (6.7 mmol) (Boc)20 was added to a stirred ice could be crystallised from methylene chloride/hexane. 20 minutes more methylene chloride was added and the carbonate solution. Drying (magnesium sulphate) and removal of the solvent in vauo left a residue which cold solution of 1.81 g (6.4 mmol) 4-The yield was 1.66 g (68%). 35

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(ii) N-{N-4-amidino benzyl}tert-butyl carbamate

(iii) N-[4-amidino cyclohexyl methyl]tert-butyl

vacuo gave 3.8 g (87%) of the title compound. chloride, drying of the combined organic phases sodium hydroxide. Subsequent extraction with methylene (potassium carbonate) and removal of the solvent in dissolved in water and the solution was made basic with the solvent was removed in vacuo. The residue was for 20 h. The catalyst was removed by filtration and the presence of 863 mg 5% rhodium on alumina at 3.4 MPa butyl carbamate was hydrogenated in 100 ml metanol in 17 mmol of the acetate of N-[4-amidino benzyl]tert-

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methyl] tert-butyl carbamate (iv) N-[N-4-(benzyloxycarbonyl)amidino cyclohexyl

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of ethyl acetate yielded 2.49 g (80%) of the title with methylene chloride containing increasing amounts applied on a column of silica and subsequent elution hydrogen carbonate solution. The organic phase was extracted with water, dilute acetic acid, and sodium mixture was diluted with methylene chloride and methylene chloride. After 10 minutes the reaction mmol) triethyl amine, and 197 mg DMAP in 40 ml amidino cyclohexyl]tert-butyl carbamate, 1.23 ml (8.8 0°C to a stirred solution of 2.04 g (8 mmol) N-[4-1.25 ml (8.8 mmol) benzyl chloroformate was added at

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compound

cyclohexane  $(H-Pac(Z) \times 2HC1)$ . (v) 4-aminomethyl-1-(N-benzyloxy carbonylamidino)-

some of the solvent in vacuo the dihydrochloride of title compound crystallised. After 10 minutes methanol was added and upon removal of methyl]tert-butyl carbamate in 40 ml ethyl acetate. Hydrogen chloride was passed through a solution of 2 g (5.1 mmol) N-[4-(benzyloxycarbonyl)amidino cyclohexyl

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piperidine (H-Pig(Z) x HCl) 4-aminomethyl-1-(N-benzyloxy carbonylamidino)

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benzyloxycarbon ylamidino) piperidine (Boc-Pig(Z)) (i) 4-(N-tert-butyloxycarbonylaminomethyl)-1-(N-

30 25 20 (100/0, 97/3, 95/5, 90/10) as eluent to yield 5.22 gchromatography using a stepwise gradient of  $ext{CH}_2 ext{Cl}_2/ ext{MeOH}$ (37%) of the title product. evaporated. The crude product was purified by flash combined organic layer was dried ( $ext{Na}_2 ext{SO}_4$ ), filtered and twice with 0.3 M  ${
m KHSO_4}$  and once with brine. The was dissolved in  $ext{CH}_2 ext{Cl}_2$ . The organic layer was washed two days. The solvent was evaporated and the residue 60-70°C for six hours and left at roomtemperature for mixed in 25 mL ethanol. The mixture was heated at mmole) of N-benzyloxycarbonyl-S-methylisothiourea was butyloxycarbonylaminomethyl) piperidine and 8.98 g (40 7.8 g (36.4 mmole) of 4-(N-tert-

carbonylamidino) piperidine (ii) H-Pig(Z) x HCl (4-aminomethyl-1-(N-benzyloxy

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mL ethyl acetate saturated with HCl(g). The mixture was 5.22 g (13.5 mmole) of Boc-Pig(Z) was dissolved in 100

allowed to stand for one hour and then evaporated. The residue was dissolved in water and washed with a mixture of diethylether and ethyl acetate. The water layer was freeze-dried to yield 4.0 g (91%) of the title compound.

<sup>1</sup>H-NYR (D<sub>2</sub>O, 300 MHz): 6 1.40-1.60 (m, 2H), 2.05 (bd, 2H), 2.19 (m, 1H), 3.07(d, 2H), 3.34(bt, 2H), 4.08 (bd, 2H), 5.40 (s, 2H), 7.5-7.63 (m, 5H)

MS m/z 291 (M<sup>+</sup>+1)

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4-Aminoethyl-1-benzyloxycarbonylamidino piperidine (H-Rig(3))

(1) 1-Benzyloxycarbonylamidino-4-hydroxyethyl piperidine

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A mixture of 6.2 g (0.028 mol) of 4-hydroxyethyl piperidine and 3.6 g (0.028 mol) of N-benzyloxycarbonyl-S-methyl isothiourea in 50 ml of acetonitrile was refluxed overnight. Evaporation and flash chromatography on silica gel with

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<sup>1</sup>H-NYR (300 NHz, CDCl<sub>3</sub>): 6 1.1-1.85 (m, 7 H), 2.83 (bt, 2 H), 4.70 (bt, 2 H), 4.18 (bd, 2 H), 5.12 (s, 2 H), 6.9-7.2 (m, 2 H), 7.2-7.5 (m, 5 H).

ethyl acetate gave 3.5 g (41%) of the title compound.

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30 (ii) 1-Benzyloxycarbonylamidino-4-mesyloxyethyl piperidine To an ice cooled solution of 3.50 g (0.0115 mol) of 1-benzyloxy-carbonylamidino-4-hydroxyethyl piperidine, 1.15 g (0.0115 mol) of triethyl amine in 40 ml of methylene chloride and 10 ml of tetrahydrofurane was added dropwise 1.30 g (0.115 mol) of mesyl chloride.

The reaction mixture was allowed to stir for 1 h. The mixture was poured into water and the organic layer was kept. The aqueous layer was extracted with methylene chloride and the combined organic layers were washed

with water, dried (Na<sub>2</sub>SO<sub>4</sub>) and evaporated. The product was used without further purification in the next step. Yield: 4.4 g (100%). 1H NMR (500 MHz, CDCl<sub>3</sub>) d 1.15-1.3 (m, 2 H), 1.65-1.8
10 (m, 5 H), 2.84 (bt, 2 H), 3.01 (s, 3 H), 4.20 (bd, 2 H), 4.27 (t, 2 H), 5.12 (s, 2 H), 7.1-7.5 (m, 7 H).

(iii) 4-Azidoethyl-1-benzyloxycarbonylamidino piperidine

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In 100 ml of dimethylformamide was dissolved 4.4 g (0.0115 mol) of crude 1-benzyloxycarbonylamidino-4-mesyloxyethyl piperidine and 4.5 g (0.069 mol) of sodium azide was added. The mixture was heated at 100°C sodium azide was added. The mixture was heated at 100°C if or 2.5 h. It was then poured into water and extracted with ethyl acetate three times. The combined organic phase was washed with water, dried (Na<sub>2</sub>SO<sub>4</sub>) and evaporated. The residue was flash chromatographed on silica gel using ethyl acetate/heptane 1/1 as eluent.

<sup>1</sup>H-NMR (500 MHz, CDCl<sub>3</sub>) & 1.20 (dq, 2H), 1.5-1.8 (m, 5 H), 2.85 (dt, 2 H), 3.35 (t, 2 H), 4.22 (bd, 2 H), 5.13 (s, 2 H), 6.9-7.2 (b, 2 H), 7.2-7.45 (m, 5 H).

Yield: 3.0 g (79%).

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(iv) 4-Aminoethyl-1-benzyloxycarbonylamidino piperidine (H-Rig(2))

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To 30 ml of water was added 0.40 g of 10% Pd/C. Sodium 35 borohydride, 1.0 g (0.031 mol), was dissolved in 30 ml of water and was added carefully to the stirred and ice-cooled slurry of Pd/C and water. 4-Azidoethyl-1-

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- 12 sec.

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without further purification. phase was washed with water, dried (Na2SO4) and methylene chloride three times. The combined organic evaporated. The product was used in the following step made alkaline with 2 M NaOH and extracted with was washed with ethyl acetate. The aqueous phase was tetrahydrofuran was evaporated and the aqueous phase the celite was rinsed with additional water. The was added. The mixture was filtered through celite and the mixture was ice-cooled again and 30 ml of 2 M HCl slurry above. After 4 h of stirring at room temperature solution was added dropwise to the ice-cooled aqueous was dissolved in 80 ml of tetrahydrofurane and this benzyloxycarbonylamidino piperidine, 2.9 g (8.8 mmol),

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4.20 (bd, 2 H), 5.12 (s, 2 H), 6.9-7.2 (b, 2 H), 7.2-7.5 (m, 5 H). (m, 1H), 1.73 (bd, 2 H), 2.72 (b, 2 H), 2.81 (dt, 2 H), <sup>1</sup>H-NMR (500 MHz, CDCl<sub>3</sub>) & 1.1-1.5 (m, 6 H), 1.55-1.65

Pyrrolidine (H-(R,8)Nig(Z)) (3RS)-1-(N-benzyloxycarbonylamidino)-3-aminomethyl 20

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## (i) (3RS)-3-hydroxymethyl pyrrolidine

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gave the product in a quantitative yield. Filtration through hyflo and evaporation of the solvent ethanol for three days completed the reduction. 0.26 MPa over 1.6 g Pd/C (10%) in 5 ml water/150 ml reaction was not completed. Continued hydrogenation at over night. After filtration through hyflo and evaporation of the solvent the  $^1\mathrm{H-NMR}$  showed that the ethanol and the mixture was hydrogenated at 0.26 MPa mixed with 1.6 g Pd/C (10%), 5 ml water and 150 ml Pyrrolidine (See H-(R,S)Hig(Z) (i) vide supra) was 16.4 g (0.0857 mole) (3RS)-1-benzyl-3-hydroxymethyl

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hydroxymethyl pyrrolidine (ii) (3RS)-1-(N-benzyloxycarbonylamidino)-3-

15 10 of the product. using  $CH_2Cl_2/MeOH$  95/5 as eluent to yield 0.70 g (25%) crude product was purified by flash chromatography water, dried  $(\mathrm{Na_2SO_4})$ , filtered and evaporated. The the mixture was dissolved in  $ext{CH}_2 ext{Cl}_2$  , washed once with temperature over night. The solvebt was evaporated and 60°C for three hours followed by stirring at room therefore dissolved in 15 ml acetonitrile and heated to the reaction was not completed. The mixture was The mixture was evaporated and the  $^1H ext{-NMR}$  showed that methylisourea was dissolved (the amine not very soluble) in toluene and heated to 60°C for three hours followed by stirring at room temperature over night. and 2.29 g (0.011 mole) N-benzyloxycarbonyl-0-1.01 g (0.01 mole) (3RS)-3-hydroxymethyl pyrrolidine

#### 20 MS m/z 278 (M+1)

mesyloxymethyl pyrrolidine (iii) (3RS)-1-(N-benzyloxycarbonylamidino)-3-

30 25 solution and extracted twice with  $\mathrm{CH_2Cl_2}$ . The combined organic layer was dried  $(Na_2SO_4)$ , filtered and The water layer was made neutral with 10 M NaOHsolution. The water layer was washed once with  $\mathrm{CH_2Cl}_2$ . ethyl acetate and extracted with a 0.3 M  $ext{KHSO}_4$ solvent was evaporated and the residue was dissolved in chloride in 3 ml diethyl ether was slowly added and the reaction mixture was stirred at 0°C for three hours. The cooled to 0°C. 0.25 ml (3.29 mmole) methanesulphonyl 15 ml diethyl ether/ $\mathrm{CH_2Cl_2}$  1/1 and the mixture was and 0.70 ml (5.05 mmole) triethylamine was dissolved in benzyloxycarbonylamidino)-3-hydroxymethyl pyrrolidine 0.7 g (2.53 mmole) (3RS)-1-(N-

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evaporated to yield 0.450 g (50%) of the title combound. (1v) (3RS)-1-(N-benzyloxycarbonylamidino)-3-azidomethyl pyrrolidine

chromatography using  ${
m CH_2Cl_2/MeOH}$  95/5 as eluent to yield benzyloxycarbonylamidino)-3-mesyloxymethyl pyrrolidine and 0.124 g (1.9 mmole) of sodium azide were dissolved in 10 ml dimethylformamide and heated to 60°C for four night. Water was added and the mixture was extracted evaporated. The crude product was purified by flash hours followed by stirring at room temperature over twice with toluene/ethyl acetate 2/1. The combined organic layer was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and 0.450 g (1.27 mmole) (3RS)-1-(N-0.262 g (68%) of the product.

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MS m/z 303 (M+1)

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(v) (3RS)-1-(N-benzyloxycarbonylamidino)-3-aminomethyl pyrrolidine (H-(R,S)Nig(Z))

stream of nitrogen was passed. 98 mg NaBH, in 2.6 ml  $\mathrm{H}_2\mathrm{O}$ pressurea and the remaining water layer was washed once mesyloxymethyl pyrrolidine dissolved in 7 ml MeOH. The mixture was allowed to stand for one hour. 5 ml 1M HCl 32 mg Pd/C (10%) and 2.6 ml  ${\rm H_2O}$  was mixed and a gentle and extracted several times with  ${
m CH}_2{
m Cl}_2$ . The combined was added and the mixture was filtered through hyflo. with ethyl acetate, made alkaline with NaOH-solution was added folowed by a slow addition of 262 mg (0.87 The organic solvent was evaporated under reduced evaporated to yield 130 mg (54%) of the product. organic layer was dried ( $\mathrm{Na_2SO_4}$ ), filtered and mmole) (3RS)-1-(N-benzyloxycarbonylamidino)-3-MS m/z 277 (M+1)

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(3R8)-1-(M-benzyloxycarbonylamidino)-3-aminosthyl pyrrolidine (H-(R,8)Hig(Z))

(i) (3RS)-1-benzyl-3-hydroxymethyl pyrrolidine

 ${\tt Na_2SO_4/cellite}$ , filtered and evaporated to give (20.3 g) water, 18 ml 3.75 M NaOH-solution and 6 ml water . The diethyl ether under an argon-atmosphere. The mixture slurry of 6.22 g lithium aluminium hydride in 160 ml was stirred over night and then heated to reflux for followed by a slow addition of, in that order, 6 ml one hour. The reaction mixture was cooled to room temperature and 0.2 g of  ${
m Na_2SO_4}$  x 10  ${
m H_2O}$  was added methoxycarbonyl pyrrolidine was slowly added to a 25.2 g (0.1063 mole) (3RS)-1-bensyl-2-oxo-4slurry was dried from excess of water with of the product. 15

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<sup>1</sup>H-NMR (CDCl<sub>3</sub>, 300 MHz): 6 1.64-1.77 (m, 1H), 1.93-2.07 1H), 2.82 (m, 1H), 3.52 (dd, 1H), 3.59 (s, 2H), 3.67 (m, 1H), 2.27-2.40 (m, 2H), 2.51 (dd, 1H), 2.62 (dd, (dd, 1H), 7.15-7.40 (m, 5H)

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(ii) (3RS)-1-benzyl-3-chloromethyl pyrrolidine 25 To a refluxing solution of 15.3 g (0.08 mole) (3RS)-1benzyl-3-hydroxymethyl pyrrolidine in 220 ml CHCl3 was 60 ml CHCl3, and the reflux was continued for one hour. slowly added a solution of 330 ml thionyl chloride in The mixture was evaporated and the residue was dissolved in water.

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made alkaline with 0.2 M NaOH-solution. The water layer combined organic layer was dried  $(Na_2SO_4)$ , filtered and The water layer was washed with ethyl acetate and then evaporated to give the product in a quantitative yield was extracted three times with ethyl acetate and the 32

(16.8 g).

1H-NMR (CDCl<sub>3</sub>, 300 MHz): & 1.55 (m, 1H), 2.05 (m, 1H), 2.38 (dd, 1H), 2.48-2.64 (m, 3H; thereof 2.58 (t, 2H))), 2.73 (dd, 1H), 3.51 (d, 2 H), 3.60 (s, 2H), 7.2-7.4 (m, 5H)

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# (iii) (3RS)-1-benzyl-3-cyanomethyl pyrrolidine

16.8 g (0.08 mole) (3RS)-1-benzyl-3-chloromethyl pyrrolidine and 5.88 g (0.12 mole) of sodium cyanide was dissolved in 250 ml dimethyl sulfoxide. The mixture was stirred at 60°C for two days and at room temperature for one week. Water was added and the mixture was extracted three times with ethyl acetate. The combined organic layer was washed with brine, dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and evaporated to yield 14.7 g (92%) of the product.

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# (iv) (3RS)-1-benzyl-3-aminoethyl pyrrolidine

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14.7 g (0.0734 mole) (3RS)-1-benzyl-3-cyanomethyl pyrrolidine dissolved in 220 ml diethyl ether was slowly added to a slurry of 2.94 g of lithium aluminium hydride in 74 ml diethyl ether under an argon atmosphere. The mixture was stirred over night, and 6 ml water, 18 ml 3.75 M NaOH-solution and 6 ml water was added to the mixture. The slurry was dried from excess of water with Na<sub>2</sub>SO<sub>4</sub>/cellite, filtered by suction and evaporated to yield 14.84 g (99%) of the product.

<sup>1</sup>H-NMR (CDCl<sub>3</sub>, 300 MHz): 6 1.41 (m, 1H), 1.51 (q, 2H), 1.90-2.10 (m, 2H; thereof 2.05 (dd, 1H))), 2.18 (m,

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1H), 2.43 (m, 1H), 2.55-2.73 (m, 3H), 2.80 (apparent t, 1H), 3.58 (apparent d, 2H), 7.15-7.4 (m, 5H)

(v) (3RS)-1-benzyl-3-(N-tert-5 butyloxycarbonylaminoethyl) pyrrolidine

To a mixture of 14.84 g (0.0726 mole) (3RS)-1-benzyl-3aminoethyl pyrrolidine, 72.6 ml IM NaOH-solution, 76 ml
water and 145 ml THF was added 17.44 g (0.08 mole) ditert-butyl dicarbonate and the mixture was stirred over
night. The solution was concentrated and extracted
three times with ethyl acetate. The combined organic
layer was washed with brine, dried (Na<sub>2</sub>SO<sub>4</sub>), filtered
and evaporated. The crude product was purified by flash
chromatography using a stepwise gradient of CH<sub>2</sub>Cl<sub>2</sub>/MeOH
( 95/5, 90/10 ) as eluent to yield 14.69 g (80%) of the
product.

1H-NMR (CDC13, 300 MHz): & 1.25-1.65 (m, 12H; thereof 20 1.40 (s, 9H)), 1.90-2.25 (m, 3H), 2.46 (m, 1H), 2.67 (m, 1H), 2.80 (apparent t, 1H), 3.09 (m, 2H), 3.59 (s, 2H), 4.60 (bs, NH), 7.15-7.35 (m, 5H)

(vi) (3RS)-3-(N-tert-butyloxycarbonylaminoethyl)
25 pyrrolidine

3.1 g (0.01 mol) (3RS)-1-benzyl-3-(N-tert-butyloxycarbonylaminoethyl) pyrrolidine was hydrogenated at 0.28 MPa over 0.6 g of Pearlman's catalyst (Pd(OH)<sub>2</sub>) in 40 ml ethanol (95%) over night. After filtration of the catalyst through cellite and evaporation of the solvent <sup>1</sup>H-NMR showed that the reaction was not completed. Therefore 0.6 g of Pearlman's catalyst was added in 40 ml ethanol (95%) once more and the mixture was treated under H<sub>2</sub>-atmosphere at 0.28 MPa over night. Filtration through

cellite and evaporation of the solvent gave the product

in a quantitative yield (2.18 g)

MS m/z 214 (M<sup>+</sup>)

(vii) (3RS)-1-(N-benzyloxycarbonylamidino)-3-aminoethyl pyrrolidine (H-(R,S)Hig(Z))

(0.0125 mole) N-benzyloxycarbonyl-S-methylisothiourea was dissolved in 30 ml toluene and heated to 60-70°C butyloxycarbonylaminoethyl) pyrrolidine and 2.81 g 2.18 g (0.0102 mmole) (3RS)-3-(N-tert-ទ

temperature for one day. 0.3 M  ${
m KHSO_4}{ ext{-}}{
m solution}$  was added which time the Boc group was removed. The acidic water phase was made alkaline and extracted four times with  $\mathrm{CH_2Cl_2}_{,\cdot}$  The combined organic layer was dried (Na\_SSO,), and the water layer was washed with a mixture of the filtered and evaporated to yield 2.0.g (51%) of the toluene and ethyl acetate and left for 2 days under for eight hours followed by stirring at room 15

H-NMR (CDCl<sub>3</sub>,330 K, 300 MHz): 6 1.45-1.7 (m, 3H), 2.07 1H), 3.33 (apparent q, 1H), 3.45-3.80 (m, 2H), 5.12 (s, (m, 1H), 2.26 (m, 1H), 2.74 (t, 2H), 3.00 (apparent t, 2H), 6.72 (bs, 2 NH), 7.15-7.45 (m, 5H)

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title compound.

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(4RS)-1,3-diaza-2-tosylimino-4-aminoethyl cyclohexane (H-(R,S)Itp(Ts)) (i) (4RS)-1,3-diaza-2-tosylimino-4-carboxy cyclohexane Prepared using the same method as described in Journal 30

of Org. Chem., p. 46, 1971.

(11) (4RS)-1,3-diaza-2-tosylimino-4-hydroxymethyl cyclohexane 32

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12.9 g (145 mmol) LiAlH, was carefully added to a cold (4RS)-1,3-dlaza-2-tosylimino-4-carboxy cyclohexane in slurry (ice bath temperature) of 9.9 g (33 mmol) of 330 mL dry THF. The reaction was stirred at room

and celite to the mixture, and filtered. Evaporation of temperature over night. The reaction mixture was worked water, 38.7 g 3.75 M NaOH, 12.9 g water,  ${\rm Na_2SO_4},~{\rm CH_2Cl_2}$ up according to Fieser & Fieser , e.g by adding 12.9 g the solvent gave 7.0 g (75%) of the desired product.

MS m/z 284 (M+ 1)

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(iii) (4RS)-1,3-diaza-2-tosylimino-4-mesyloxymethyl cyclohexane

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organic phase was separated, dried( ${
m Na}_2{
m SO}_4$ ), filtered and cyclohexane in 6.9 mL (49.4 mmol) triethylamine and 125 evaporated to give the title compound in quantitative 2.9 mL MsCl (37.1 mmol) was added carefully to a cold (ice bath temperature) slurry of 7.0 g (24.7 mmol) of  $^{\mathrm{mL}}$  CH<sub>2</sub>Cl<sub>2</sub>. Water was added after 1h 15 min and the (4RS)-1,3-diaza-2-tosylimino-4-hydroxymethyl 20

MS m/z 362 (M+ +1)) 25

yield.

(iv) (4RS)-1,3-diaza-2-tosylimino-4-cyanomethyl cyclohexane

was dissolved in 75 mL DMSO. After stirring at 40°C for 60 hours an additional amount of 0.31 g (6 mmol) NaCN precipitated out of the solution. They where filtered mesyloxymethyl cyclohexane and 1.3 g (27.2 mmol) NaCN 8.9 g (24.7 mmol) of (4RS)-1,3-diaza-2-tosylimino-4was added and the solution was stirred at 65°C for three hours. 150 mL water was added and crystals off and dried to give 5.4 g (75%) of the desired 32 30

product.

MS m/z 293 (M+ + 1)

(4RS)-1,3-diaza-2-tosylimino-4-aminoethyl cyclohexane (H-(R,S)Itp(Ts))

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mixture was filtered and the solvent removed in vacuo THF. After stirring for 2 hours 1 g  $H_2O$ , 3 g 3.75M to give 2.2 g (90%) of the desired product. NaOH, 1 g H<sub>2</sub>O, Na<sub>2</sub>SO<sub>4</sub>, celite and CH<sub>2</sub>Cl<sub>2</sub> was added. The diaza-2-tosylimino-4-cyanomethyl cyclohexane in 90 mL temperature) slurry of 2.4 g (8.2 mmol) of (4RS)-1,3-935 mg  $LiAlH_4$  was added carefully to a cooled (ice bath

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3.44-3.53 (m, 1H), 7.28 (d, 2H), 7.71 (d, 2H) 2H, partially overlapping with the solventsignal), <sup>1</sup>H NMR (500 MHz, MeOD); & 1.52-1.71 (m, 3H), 1.88-1.96 (m, 1H), 2.37 (s, 3H), 2.64-2.73 (m, 2H), 3.2-3.4 (m,

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(H-(8) Itp(Ts)) (48)-1,3-diaza-2-tosylimino-4-aminoethyl cyclohexane 20

diaminobutyric acid. (R,S)Itp(Ts) starting from optically pure 2,4-Prepared in the same way as described for H-

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2.68-2.82 (m, 1H), 2.86-2.98 (m, 1H), 3.22-3.44 (m, 2H), 3.45-3.58 (m, 1H), 7.19 (d, 2H), 7.72 (d, 2H) 1.48-1.69 (m, 3H), 1.84-1.95 (m, 1H), 2.37 (s, 3H), <sup>1</sup>H NMR (300.13 MHz, CDCl<sub>3</sub>); & 0.97-1.15 (s broad, 1H),

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13C NMR (300.13 MHz, CDCl<sub>3</sub>); & guanidinecarbon 154.05

### H-Aze-OEt x HCl

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Prepared in the same way as described for H-Pic-OEt x

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HCl from H-Aze-OH (vide infra).

### H-Aze-OMe x HC]

ហ Seebach D. et. al.in Liebigs Ann. Chem., p. 687, 1990. Prepared according to the procedure described by

### H-Pab(Z) x HC1

15 10 to give the title compound in almost quantitative precipitates out of the solution. After filtration the yield. precipitate was washed 2 times with cold EtOH and dried 1 g/10 ml) where upon H-Pab(Z) x HCl immedeately propanol to a solution of crude H-Pab(Z) in EtOH (about Prepared by adding 1 mole equivalent of 5 M HCl in iso-

### H-Pic-OEt x HC1

- 25 20 removed and the mixture was refluxed for 2.5 h. The was added dropwise over 15 min. The ice bath was solvent was evaporated and the product was obtained as was cooled in an ice bath and 17 ml of thionyl chloride L-Pipecolinic acid, 4.0 g (0.031 mol), was slurried in its hydrochloride salt in a quantitative yield. bubbled through until a clear solution was obtained. It 100 ml of abs. ethanol and HCl (g) was carefully
- 30 2.3-2.5 (m, 1H), 3.1-3.3 (m, 1H), 3.5-3.7 (m, 1H), 4.14 <sup>1</sup>H-NMR (300 MHz, D<sub>2</sub>O): & 1.33 (t, 3H), 1.8-2.1 (m, 5H), (dd, 1H), 4.44 (q, 2H).

## H-(R,S)betaPic-OMe x HCl

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methanol was cooled in an ice-bath and 2.76 g (23.2 A mixture of 2.0 g (15.5 mmol) nipecotic acid in 8 ml

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quantitative yield.

Boc- (R) Cha-08u

Boc-(R)Cha-OH (1 eq.), HOSu (1.1 eq) and DCC or CME-CDI dried in vacuo. (When CME-CDI was used in the reaction night. The precipitate formed during the reaction was water and dried). Evaporation of the solvent gave the filtered off, the solvent evaporated and the product dissolved in EtOAc and the organic phase washed with (1.1 eq) were dissolved in acetonitrile (about 2.5 ml/mmol acid) and stirred at room temperature over the residue, after evaporation of the  $CH_3CN$ , was title compound.

0.85-1.1 (m, 2H), 1.1-1.48 (m, 4H), 1.5-1.98 (m, 16H; thereof 1.55 (bs, 9H)), 2.82 (bs, 4H), 4.72 (bs, 1H, <sup>1</sup>H-NMR (500 MHz, CDCl<sub>3</sub>, 2 rotamers ca: 1:1 ratio) major rotamer), 4.85 (bs, 1H, minor).

Boc-(R) Hoc-OH

mixture was stirred under a hydrogen atmosphere at 0.41 hyflo and the solvent evaporated giving the product in MPa for 18 h. The catalyst was filtered off through dissolved in methanol (75 ml). Rhodium on activated aluminium oxide  $(\mathrm{Rh}/\mathrm{Al}_2\mathrm{O}_3)$ , 0,5 g was added and the Boc-(R)Hop-OH (See above), 3.2 g (11.46 mmol) was almost quantitative yield.

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H-NMR (500 MHz, CDCl3): 6 0.90 (m, 2H), 1.08-1.33 (m, 6H), 1.43 (s, 9H), 1.60-1.74 (m, 6H), 1.88 (bs, 1H), 4.27 (bs, 1H).

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Boc- (R) Hoc-08u 35

Prepared in the same way as described for Boc-(R)Cha-

was evaporated and the residue was dissolved in a small

stirred at room temperature for 20 hours. The solvent

mmol) thionyl chloride was added. The mixture was

(R,S) betapic-OMe x HCl precipitated as white crystals.

amount of methanol, diethylether was added and H-

The crystals 2.57 g (92%) were isolated by filtration.

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with water, dried (Na $_2$ SO $_4$ ) and evaporated to give 28.3 g of the title compound. The crude material was dissolved showed the presence of about 25 % of the methyl ester in 500 ml of THF and 300 ml of water and 20 g of LiOH were added. The mixture was stirred overnight and the ethyl acetate. The combined organic layer was washed acidified with  $KHSO_4$  and extracted three times with THF was evaporated. The remaining water phase was

1H-NMR (300 MHz, CDCl3): 6 0.9-1.7 (m, 20H), 4.0-4.2 (m, (83 %) of the desired product.

1H), 5.2 (d, 1H).

Boc-(R) cg1-08u

Evaporation of the solvent gave the title compound in stirring for 3 days the precipitated DCU was filtered reaction was allowed to reach room temperature. After To an ice-cold solution of 2.01 g (7.81 mmol) of Boc-(R)Cgl-OH and 1.83 g (15.6 mmol) of HOSu in 25 ml of dissolved in EtOAc and the organic phase was washed with  $\rm H_2O$ , KHSO4, NaHCO3, brine and dried(Na<sub>2</sub>SO<sub>4)</sub>.  $\mathrm{CH_3CN}$  was added 1.69 g (8.2 mmol) of DCC and the of f and the solvent evaporated. The residue was 35 ဗ္ဗ

Boc-(R)-Pg1-OH, 32.6 g (0.13 mol), was dissolved in 300 ml of methanol and 5 g of  $\mathrm{Rh}/\mathrm{Al}_2\mathrm{O}_3$  was added. The ព

Boc- (R) Cg1-OH

solution was hydrogenated at 5.2 to 2.8 MPa for 3 days. After filtration and evaporation of the solvent NMR

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OSu from Boc-(R)Hoc-OH.

### Boc-(R) Pro(3-(S) Ph) -OH

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Prepared according to the method described by J.Y.L Chung et al in Journal of Organic Chemistry, No 1, pp. 270-275, 1990.

### Boc-(R)Cgl-Aze-OH

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### (i) Boc-(R)Cgl-Aze-OMe

To a stirred mixture of 3.86 g (15 mmol) Boc-(R)Cgl-OH, 2.27 g (15 mmol) H-Aze-OMe x HCl and 2.75 g (22.5 mmol) DMAP in 40 mL CH<sub>3</sub>CN at 5°C was added 3.16 g (16.5 mmol) EDC. The reaction mixture was stirred at room temperature for 48h. The solvent was evaporated and the residue was dissolved in 150 ml EtoAc and 20 ml H<sub>2</sub>O. The separated organic layer was washed with 2 x 20 ml 0.5 M KHSO<sub>4</sub>, 2 x 10 ml NaHCO<sub>3</sub>(saturated), 1 x 10 ml H<sub>2</sub>O, 1 x 10 ml brine and dried (MgSO<sub>4</sub>). Evaporation of the solvent gave 4.91 g (92 %) of the title compound which was used without further purification in the next step.

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### (ii) Boc-(R)Cgl-Aze-OH

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The hydrolysis of Boc-(R)Cgl-Aze-OMe was carried out according to the procedure described for Boc-(R)Cha-

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Pic-OEt (vide infra). The product was crystallized from EtOH/acetone/water (1/1/3.95) yield 80 %.

1H-NMR (500 MHz, CDCl<sub>3</sub>): 6 0.85-1.3 (m, 5H), 1.40 (s, 9H), 1.5-1.9 (m, 6H), 1.95-2.2 (m, 2H), 3.92 (m, 1H), 4.09 (m, 1H), 4.35 (m, 1H), 4.95 (m, 1H), 5.16 (bd, 1H).

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### Boc-(R)Cgl-Pic-OH

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### (i) Boc-(R)Cgl-Pic-OMe

Pivaloyl chloride (1.0 ml, 8.1 mmol) was added to a solution of Boc-(R)Cgl-OH (2.086 g, 8.1 mmol) and triethyl amine (1.13 ml, 8.1 mmol) in toluene (25 ml) and DMF (5 ml). A mixture of H-Pic-OMe x HCl (1.46 g, 8.1 mmol) and triethyl amine (1.13 ml, 8.1 mmol) in DMF (20 ml) was subsequently added at ice bath temperature. The reaction mixture was slowly allowed to warm up to room temperature and after 24 h it was diluted with water and extracted with toluene. After washing with 0.3 M KHSO<sub>4</sub>, 10% Na<sub>2</sub>CO<sub>3</sub> and brine the solvent was removed in vacuo to give 2.52 g (81%) of colorless oil which was used without further purification.

1H-NMR (500 MHz, CDCl<sub>3</sub>, 2 rotamers, 5:1 ratio) & 0.8-1.8
(m, 25H), 2.25 (d, 1H), 2.75 (t, 1H, minor rotamer),
3.3 (t, 1H), 3.7 (s, 3H), 3.85 (d, 1H), 4.3 (t, 1H,
minor rotamer), 4.5-4.6 (m, 1H), 5.25 (d, 1H), 5.30 (d,
1H).

### (ii) Boc-(R)Cgl-Pic-OH

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Prepared according to the procedure for hydrolysis of Boc-(R)Cha-Pic-OEt (vide infra) using the product from (1) above. The product was crystallized from disopropyl ether and hexane.

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<sup>1</sup>H-NMR (500 MHz, CDCl<sub>3</sub>, 2 rotamers, 5:1 ratio) 6 0.8-1.8 (m, 25H), 2.3 (d, 1H), 2.8 (t, 1H, minor rotamer), 3.3 1H), 5.1 (s, 1H, minor rotamer), 5.3 (d, 1H), 5.40 (d, (t, 1H), 3.9 (d, 1H), 4.4 (t, 1H, minor), 4.5-4.6 (m, 8

### Boc-(R) Cgl-Pro-OH

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water and 1.18 g (29.7 mmol) of sodium hydroxide. 2.8 g (7.8 mmol ) of Boc-(R) Cgl-OSu in 10 ml DMF was added to The organic phase was washed once with water and once washed with ethyl acetate, acidified with 0.3 M  $\rm KHSO_4^$ solution and extracted three times with ethyl acetate. with brine, dried  $(\mathrm{Na}_2\mathrm{SO}_4)$ , filtered and evaporated to 3.59 g (31.24 mmol) of L-proline was mixed with 20 ml mixture was stirred for three days. The solvent was evaporated and water was added. The water phase was additional 30 ml of DMF was added and the reaction the mixture . Because of solubility problem an yield 2.3 g (83 %) of the product.

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'H-NMR (300 MHz, CDCl<sub>3</sub>): 6 0.89-2.17 (m, 23H), 2.37 (m, 1H), 3.55 (q, 1H), 3.90 (bs, 1H), 4.28 (t, 1H), 4.52 (bs, 1H), 5.22 (bs, 1H (NH)).

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Boc-(R)Cha-Aze-OH

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Pic-OH starting from Boc-(R)Cha-OH and H-Aze-OEt x HCl Prepared in the same way as described for Boc-(R)Cha-

(vide infra). 30

### Boc-(R) Cha-Pro-OH

hydroxide (750 ml). Boc-(R)Cha-OSu (170 mmol) dissolved H-(5)Pro-OH (680 mmol) was dissolved in 0.87 M sodium in DMF (375 ml) was added dropwise during 20 min. The reaction mixture was stirred at room temperature for

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finally the product dried in vacuo to yield Boc-(R)Chaand once with brine. After drying over sodium sulphate dissolved in diethyl ether, the solvent evaporated and layers were combined and washed three times with water Pro-OH as a white powder in almost quantitative yield. extracted three times with ethyl acetate. The organic and evaporation of the solvent, the syrupy oil was 20 h. The mixture was acidified (2M KHSO $_4$ ) and

3.48 (m, 1H), 3.89 (m, 1H), 4.55 (m, 2H), 5.06 (m, 1H); 9H)), 1.55-1.8 (m, 5H), 1.8-2.15 (m, 3H), 2.47 (m, 1H),  $^{1}\mathrm{H-NMR}$  (500 MHz, CDCl $_{3}$ ,mixture of two rotamers 9:1)  $^{6}$ Resolved signals from the minor rotamer appears at d 0.8-1.05 (m, 2H), 1.05-1-55 (m, 15H; thereof 1.5 (bs, 2.27 (m), 3.58 (m), 4.33 (m), 5.0 (m). ដ

## Boc-(Me) (R) Cha-Pro-OSu

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## (i) Boc-(Me) (R) Cha-Pro-OH

(R)Cha-Pro-OH starting from Boc-(Me) (R)Cha-OSu and H-Prepared in the same way as described above for Boc-Pro-OH. 2

(ii) Boc-(Me)(R)Cha-Pro-OSu 25

Prepared in the same way as described for Boc-(R)Cha-OSu starting from Boc-(Me)(R)Cha-Pro-OH.

### Boc-(R)Cha-Pic-OH ဗ္ဗ

## (ia) Boc-(R)Cha-Pic-OEt

11.2 g (0.0265 mol) of CME-CDI were added. The ice bath Boc-(R)Cha-OH, 6.3 g (0.023 mol), was dissolved in 150 ml of CH2Cl2. The solution was cooled in an ice bath and 6.3 g (0.047 mol) of N-hydroxybenzotriazole and

dilute  $\mathrm{KHSO_4}$  (aq),  $\mathrm{NaHCO_3}$  (aq) and water. The organic further purification. (89 %) of Boc-(R)Cha-Pic-OEt which was used without layer was dried  $(Na_2SO_4)$  and evaporated to give 7.7 g residue was dissolved in ethyl acetate and washed with stirred for 3 days. The solvent was evaporated and the removed after 15 min and the reaction mixture was by addition of N-methylmorpholine. The ice bath was mol) was added and the pH adjusted to approximately 9 and cooled in an ice bath. H-Pic-OETXHC1, 4.1 g (0.021 evaporated and the residue dissolved in 150  $\mathtt{ml}$  of DMF stirred for 4 h at room temperature. The solvent was was removed after 15 min and the reaction mixture was

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20 15 minor), 4.77 (bq, 1H, major), 4.90 (bd, 1H, minor), 5.28 (bd, 1H, major), 5.33 (bd,1H, major). (bd, 1H, major), 4.15-4.3 (m, 2H), 4.5-4.7 (m, 2H, 1H), 2.88 (bt, 1H, minor), 3.30 (bt, 1H, major), 3.80 1.45 (bs, 9H), 2.01 (bd, 1H, major rotamer), 2.31 (bd, 1.0 (m, 2H), 1.1-1.9 (m, 29H; thereof 1.28 (t, 3H)),  $^{1}\text{H-NMR}$  (500 MHz, CDCl $_{3}$ , 2 rotamers, 3:1 ratio ) d 0.7-

### (ib) Boc-(R)Cha-Pic-OMe

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removal of the solvent in vacuo gave 1.16 g of the 0.3 M KHSO4, 10%  $\mathrm{Na_2CO_3}$  and brine. Drying (  $\mathrm{MgSo_4}$ ) and title compound. reaction mixture and the organic phase was washed with another 18 h. Water and toluene was added to the triethyl amine was added and stirring was continued for slurry after 45 minutes. After 2 h 100  $\mu$ l (0.72 mmol) triethyl amine in 5 ml DMF was added to the resulting pipecolate hydrochloride and 463  $\mu$ l (3.32 mmol) 2 ml DMF. A mixture of 596 mg (3.32 mmol) methyl (S)stirred mixture of 875 mg (3.22 mmol) Boc-(R)Cha-OH and 450  $\mu l$  (3.23 mmol) triethyl amine in 10 ml toluene and 400  $\mu l$  (3.23 mmol) of pivaloy1 chloride was added to a

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## (ii) Boc-(R)Cha-Pic-OH

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crystallized from diisopropyl ether/hexane. without further purification. The compound can be 4.9 g (94 %) of Boc-(R)Cha-Pic-OH which was used washed with water, dried ( $\mathrm{Na_2SO_4}$ ) and evaporated to give acidified with  $ext{KHSO}_4$  (ag) and extracted three times with ethyl acetate. The combined organic phase was THF was evaporated and the aqueous solution was mixture was stirred at room temperature overnight. The 100 ml of THF, 100 ml of water and 7 g of LiOH. The Boc-(R) Cha-Pic-OEt, 5.6 g (0.014 mol), was mixed with

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15 the ethyl ester in (ii). hydrolysed using the same procedure as described for The methyl ester formed in procedure (ib) above can be

25 20 major rotamer), 1.46 (s, 9H, minor)), 2.33 (bd, 1H), 1H, major), 5.56 (m, 1H, major). 4.77 (bq, 1H, major), 5.03 (bs, 1H, minor), 5.33 (bd) 1H, major), 4.57 (bd, 1H, minor), 4.68 (m, 1H, minor), 2.80 (bt, 1H, minor), 3.33 (bt, 1H, major), 3.85 (bd, 1.1 (m, 2H), 1.1-2.1 (m, 27H; thereof 1.43 (s, 9H, 1H-NMR (500 MHz, CDCl<sub>3</sub>, 2 rotamers, 3.5:1 ratio) & 0.8-

## Boc-(R)Cha-(R,S)betapic-OH

## (i) Boc-(R)Cha-(R,S)betaPic-OMe

35 30 methyl morpholine was added and the reaction mixture was stirred for 24 h. The solvent was evaporated and H-(R,S) betaPic-OMe x HCl and 1.62 ml (14.6 mmol) 4-Nthe residue was dissolved in toluene and some After stirring for 1 h and 30 minutes 1.3 g (7.3 mmol) (7.3 mmol) 4-N-methyl morpholin in 20 ml acetonitrile. solution of 2.0 g (7.3 mmol) Boc-(R)Cha-OH and 0.81 ml Pivaloyl chloride 0.9 ml (7.3 mmol) was added to a

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solution, and drying with  ${\sf Na}_2{\sf SO}_4$  the solvent was removed acetate (7/3) as eluent gave 2.4 g (83\$) of the desired diethyleter. After washing with 0.3 M  ${
m KHSO_4}$  and  ${
m KHCO_3-}$ in vacuo. Flash chromatography using heptane/ethyl

product.

## (ii) Boc-(R)Cha-(R,S)betaPic-OH

acetate, dried over  $\mathrm{Na_2SO_4}$  and evaporated to give 2.0 g of LiOH in 35 ml water was added. After stirring for 5 (R,S)betaPic-OMe was dissolved in 35 ml THF and 2.1 g h the THF was removed in vacuo. The aqueous phase was At room temperature 2.35 g (5.9 mmol) of Boc-(R)Chaacidified with 2M  ${
m KHSO_4}$  and extracted with ethyl (89%) of the product.

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Boc-(R) Cha-Val-OH

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### (i) Boc-(R)Cha-Val-OMe

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removed in vacuo and ether and toluene was added to the Boc-(R)Cha-OH and 3.5 ml (25 mmol) triethyl amine in 50 3.1 ml (25 mmol) pivaloyl chloride was added at ambient followed by drying (MgSO $_{\phi}$ ) and removal of the solvent chromatography using toluene/ethyl acetate as eluent. in vacuo gave a residue which was subjected to flash ml DMF. After 3 hours 4.16 g (25 mmol) valine methyl crystals of DMAP were added and the reaction mixture ester hydrochloride in 50 ml DMF and 3.5 ml triethyl temperature to a stirred mixture of 6.75 g (25 mmol) amine was added. After stirring over night , a few was heated to 50°C for 5 minutes. The solvent was The yield of the title compound was 6.99 g (73%). residue. Washing with 0.3 M KHSO4 and 10%  ${\rm Na_2CO_3}$ 

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(ii) Boc-(R) Cha-Val-OH

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remoyed in vacuo and the remaining solution was diluted 5.6 g (230 mmol) lithium hydroxide in 75 ml THF and 75 followed by drying (MgSO $_4$ ) and removal of the solvent A mixture of 8.73 g (23 mmol) Boc-(R)Cha-Val-OMe and with water and extracted with ether. Acidification in vacuo gave 8.15 g (96%) of the title compound. ml of water was stirred for 4 hours. The THF was with 2 M  $KHSO_4$  and extraction with ethyl acetate

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### (i) Boc-(R)Hoc-Aze-OEt

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temperature. The solvent was evaporated and the residue The solution was cooled in an ice bath and 0.77 g (4.0 diluted  ${
m MHCO}_3$ , brine, dried with  ${
m MaSO}_4$  and evaporated. At room temperature 1.0 g (3.5 mmol) Boc-(R)Hoc-OH and dissolved in 20 ml DMF. 0.58 g (3.5 mmol) H-(R)Aze-OH addition of N-methyl morpholin. The reaction mixture Flash chromatography (1% EtOH in  ${
m CH}_2{
m Cl}_2$  and heptane: 0.95 g (7.0 mmol) HOBt was dissolved in 15 ml  ${\rm CH_2Cl_2}$ . mmol) of EDC was added. The ice bath was removed and was added and the pH adjusted to approximately 9 by partitioned between water and toluene. The organic was stirred for one day. The reaction mixture was the reaction mixture was stirred for 3 h at room Etoic) gave 0.35 g (25%) of the desired product. phase was separated and washed with 0.3 M KHSO $_4$ , 25

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### (ii) Boc-(R)Hoc-Aze-OH 30

OEt was dissolved in 10 ml THF and 0.59 g of LiOH in 10 and extracted with ethyl acetate, dried over  ${\tt Na_2SO_4}$  and  ${\tt KHSO_4}$  was added and the THP was removed in vacuo. The agueous phase was then made acidic with more 2M  $_{\rm KHSO_4}$ At room temperature 0.65 g (1.6 mmol) Boc-(R)Hoc-Azeml water was added. After stirring for 24 hours 2 M  $\,$ 

evaporated to give 0.5 g (85%) of the title compound.

### Boc-(R)Hoc-Pro-OB

Pro-OH from Boc-(R)Hoc-OSu. Prepared in the same way as described for Boc-(R)Cha-

a minor rotamer appears at: 6 1.92, 2.25, 3.58, 4.20 4.43 (m, 1H), 4.52 (bd, 1H), 5.26 (bd, 1H), signals of 2.14 (m, 2H), 2.34 (m, 1H), 3.48 (m, 1H), 3.85 (m, 1H), (m, 7H), 1.36-1.48 (bs, 9H), 1.48-1.78 (m, 7H), 1.98-1H-NMR (500 MHz, CDC13): 6 0.80-0.94 (m, 2H), 1.05-1.36

#### 15 Boc-(R)Hoc-Pic-OH

### (i) Boc-(R)Hoc-Pic-OMe

20 Oft (vide supra) from Boc-(R)Hoc-OH and H-Pic-OMe xPrepared the same way as described for Boc-(R)Cha-Pic-

### (ii) Boc-(R)Hoc-Pic-OH

25 Pic-OH (vide supra) from Boc-(R)Hoc-Pic-OMe. Prepared in the same way as described for Boc-(R)Cha-

rotamer appear at: 6 1.88, 2.80, 4.25, 4.55 and 4.97. 5.33 (bs, 1H), 5.44 (bd, 1H), signals of a minor (bd, 1H) 3.28 (bt. 1H), 3.85 (bd, 1H) 4,63 (m, 1H),  $^{1}\text{H-NMR}$  (500 MHz, CDCl<sub>3</sub>): 6 0.82-0.97 (m, 2H), 1.10-1.36 7H), 1.36-1.50 (bs, 9H), 1.50-1.82 (m, 11H), 2.35

### Boc-(R)Pro-Phe-OH

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### (i) Boc-(R)Pro-Phe-OMe

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H imes 50 ml water and dried (Na $_2$ SO $_4$ ). Evaporation of the which was used in the next step without further solvent gave the title compound in quantitative yield organic phase was washed with 3  $\times$  50 ml 0.3 M KHSO4, 1 reaction mixture was diluted with toluene and the was stirred over night at room temperature. The triethyl amine in 40 ml DMF was added and the reaction mixture of 2.0 g (9.29 mmol) H-Phe-OMe and 0.94 g temperature. The reaction was cooled to 0°C and a the reaction was stirred for 30 minutes at room 0.94 g (9.29 mmol) triethyl amine in 70 ml toluene/DMF (5/2) was added 1.12 g (9.29 mmol) pivaloyIchloride and To a solution of 2.0 g (9.29 mmol) Boc-(R)Pro-OH and

### (ii) Boc-(R)Pro-Phe-OH

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25 20 crystallization from diisopropyl ether to give 2.329 g solvent gave a residue which was purified by and dried  $(\mathrm{Na_2SO_4})$ . Filtration and evaporation of the (60 %) of the title compound as a white crystalline EtOAc. The combined organic phase was washed with water acidic with 1 M KHSO $_4$  and extracted with 3 x 75 ml The THF was evaporated and the water phase was made was stirred vigorously over night at room temperature. 8.93 g (21.3 mmol) LiOH x  $H_2O$  in 140 ml water/THF (1/1) A mixture of 4.0 g (10.6 mmol) Boc-(R)Pro-Phe-OMe and

### 30 Boc-(R) Pro(3-(8) Ph)-Pro-OH

## (i) Boc-(R)Pro(3-(S)Ph)-Pro-OBn

မ္ the reaction mixture was stirred for three days. The 0.84 mL NVM and 2.92 g CME-CDI at room temperature and H-Pro-OBn x HCl and 0.75 g HOBt in 11 mL DMF was added To a mixture of 1.61 g Boc-(R)Pro(3-(S)Ph)-OH, 1.65 g

100 ml  $\mathrm{H}_2\mathrm{O}$  and dried (MgSO $_4$ ). Evaporation of the solvent 300 mL BtOAc. The organic phase was washed with 2 imes 100 solvent was evaporated and the residue was dissolved in mL  $_{2}$ o, 2 x 100 mL 1 M KHSO $_{4}$ , 3 x 100 mL 1 M NaOH, 3 x chromatography using CH2Cl2/MeOH (97/1) as eluent to gave 2.53 g of an oil which was purified by flash give 2.11 g (88%) of the title compound.

## (ii) Boc-(R)Pro(3-(S)Ph)-Pro-OH

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0.94 g of Boc-(R)Pro(3-(S)Ph)-Pro-OBn was dissolved in 3.5 hours. Filtration of the catalyst and evaporation 70 ml EtOH and hydrogenated over 0.42 g 5 % Pd/C for of the solvent gave the title compound as white crystals in a quantitative yield. 12

### Boc-(R)Tic-Pro-OH

Prepared according to the procedure described by P.D. Gesellchen and R.T. Shuman in EP-0,479,489-A2.

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### BROOC-CH2-NH-CO-CH2-BI

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and dicyclohexyl carbodilmide (5 mmol). The mixture was triethyl amine (5 mmol) in 10 ml of  $\mathrm{CH}_2\mathrm{Cl}_2$  was added 2chromatography ( $\mathrm{CH_2Cl_2/MeOH,~95/5}$ ) gave a quantitative bromoacetic acid (5 mmol) dissolved in 10 ml of  $\mathrm{CH}_2\mathrm{Cl}_2$ stirred at room temperature over night and filtered. The organic phase was washed twice with 0.2 M  ${
m KHSO_4}$ , 0.2 M NaOH, brine and dried. Evaporation and flash To a solution of p-TsOH x H-Gly-OBn (5 mmol) and yield of the desired compound.

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<sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>): 6 3.89 (s, 2H), 4.05-4.11 (d, 2H), 5.19 (s, 2H), 7.06 (bs, 1H), 7.3-7.4 (m, 5H). 35

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### Boc-(R) cgl-11e-0H

Prepared in the same way as described for Boc-(R)Cgl-Pro-OH using H-Ile-OH, instead of H-Pro-OH, in 91 %

yield.

### Boc-(R) Phe-Phe-OH

### (i) Boc-(R)Phe-Phe-OMe

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under reduced pressure and the residue was dissolved in evaporation of the solvent yielded 7.5 g of Boc-(R)Phe-50 mL of ethylacetate. Extraction of the organic phase -ethylcarbodiimide hydrochloride (24.5 mmol) was added. with 50 mL aliquats of 0.5 M potassiumhydrogensulfate, Phe-OME (94%) which was used in the next step without 30 mL of acetonitrile. The solution was cooled to ice-The cooling bath was removed and the reaction mixture 4-dimethylaminopyridine (37.7 mmol) were dissolved in was stirred over night. The solvent was then removed 1 M sodiumbicarbonate and finally water followed by water temperature and 1-(3-dimethylaminopropyl)-3-Boc-(R) Phe-OH (18.8 mmol; purchased from Bachem Feinchemicalien AG), Phe-OMe (20.7 mmol) and further purification.

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### (ii) Boc-(R)Phe-Phe-OH

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pressure yielding 8.0 g of Boc-(R)Phe-Phe-OH (quant) as solvent was removed under reduced pressure. The residue with 50 mL of 0.5 M potassiumsulfate followed by 50 mL reaction mixture was stirred for 3.5 h after which the Boc-(R)Phe-Phe-OMe (16.4 mmol) was dissolved in 40 mL was dissolved in 50 mL of ethylacetate and extracted of tetrahydrofuran and lithiumhydroxide (32.8 mmol) dissolved in 20 mL of water was added rapidly. The of water. The solvent was removed under reduced

an amorphous solid.  $^1\mathrm{H}$  NMR (200 MHz, d-CHCl3); 6 7.4-6.7 (m, 10H), 5.7-4.2 (m, 6H), 1.34 (s, 9H).

### HO-CH2-COOBI

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A. et al in Bull. Soc. Chim. France., No11, pp 4018-23, Prepared according to the procedure described by Lattes

## No2) Ph-so2-ocH2-coosn Benzyl-2-(ortho-nitrobenzenesulfonyloxy)acetate (2-

give 1.18 g (34 %) of the title compound. chromatography, using heptan:EtOAc 2:1 as eluent to 3.34 g of a residue that was subjected to flash dried  $(Na_2SO_4)$ , filtered and evaporated in vacuo to give phase was washed with 20 ml 1 M HCl and 20 ml H20, were added. The phases were separated and the organic 0°C for 50 minutes and then 20 ml water and 30 ml  $\mathrm{CH_2Cl_2}$ portions during 15 minutes. The slurry was stirred at nitrobenzenesulfonylchloride was added in small 0°C and 2.8 ml (20 mmol) triethylamin was added. While  $\mathrm{CH_2Cl_2}$  and 25 ml diethylether. The mixture was cooled to keeping the temperature at 0°C 2.44 g (11 mmol) orto-1.66 g (10 mmol) benzylglykolate was dissolved in 25 ml

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7.83 (m, 5H), 7.76 (m, 3H), 8.16(dd, 1H).  $^{1}$ H-NMR (300MHz, CDCl $_{3}$ ): & 4.92 (s, 2H), 5.17 (s, 2H), 25

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Prepared according to the same procedure as described NO2) Ph-SO2-OCH2-COOBn Benzyl-2-(para-nitrobenzenesulfonyloxy)acetate (4-

above. The final compound was obtained in a crystalline to use without further purification (64 % yield). form after evaporation of the solvent and pure enough for Benzyl-2-(ortho-nitrobenzenesulfonyloxy)acetate

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7.2-7.4 (m, 5H), 8.10 (d, 2H), 8.30 (d, 2H).  $^{1}\text{H-NMR}$  (300 MHz, CDCl $_{3}$ ): & 4.79 (s, 2H), 5.13 (s, 2H),

#### Tfo-CH2COOMe

vacuo gave 9.94 g (90 %) of the title compound. minutes, and thereafter stirred at 0°C for 1 H. After washing with 0.3 M KHSO $_{\rm 4}$  and saturated NA $_{\rm 2}^{\rm CO}_{\rm 3}$ , drying pyridin in CH2Cl2 (totally 62.5 ml) at 0°C during 25 4.05 ml (50 mmol) methylglycolate and 4.04 ml (50 mmol)  $(Na_2SO_4)$  and filtration, evaporation of the solvent in dissolved in  $ext{CH}_2 ext{Cl}_2$  was added dropwise to a mixture of 10.09 ml (60 mmol) trifluorometansulfonic anhydrid

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#### 15 Tfo-CH2COOEt

Prepared in the same way as described for TfO-CH2COOMe starting with ethylglycolate.

#### 20 Tfo-CH2COO"Bu

starting with butylglycolate. Prepared in the same way as described for TfO-CH2COOMe

Tfo-CH2COOBD

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starting with HO-CH2COOBn Prepared in the same way as described for TfO-CH2COOMe

### Tfo-CH2COOTHex

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### (i) HO-CH<sub>2</sub>COO<sup>n</sup>Hex

35 (2.82 mmol) DBU. After stirring over night and reflux was added 719 mg (3.39 mmol) 1-hexyl iodide and 429 mg To 215 mg (2.82 mmol) glycolic acid in 12.8 ml  $_{\mathrm{CH_3CN}}$  - Lewis Control of the Control

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for 4 h, the solvent was evaporated, ethylacetat and 1 filtered and evaporated in vacuo to give 333 mg (74 %) M  $\mathrm{KHSO}_{4}$  was added and the phases were separated. The organic layer was washed with brine, dried (MgSO $_4$ ), of the product.

(11) TfO-CH2COO<sup>n</sup>Hex

Prepared in the same way as described for TfO-CH $_2$ COOMe starting with HO-CH2COO<sup>n</sup>Hex.

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H-Mig(Z) (3-aminomethyl-1-(N-benzyloxycarbonylamidino) azetidine (1) 3-aminomethyl-1-benzhydryl azetidine was prepared according to the literature, see A.G. Anderson, Jr., and R. Lok, J.Org.Chem., 37, 3953, 1972. 12

(11) 3-(N-text-butyloxycarbonylaminomethyl)-1-

benzhydryl azetidine 20

removed after a few minutes and the mixture was stirred mixture was cooled to 0°C and 3.03 g (13.9 mmol) of ditert-butyl dicarbonate was added. The cooling bath was filtered. Evaporation of the solvent gave 4.6 g (94 \$) at roomtemperature over night. The THF was evaporated and the residue was extracted with 3x45 mL of diethyl azetidine dissolved in 45 mL THF was added a solution of 0.56 g (13.9 mmol) NaOH in 45 mL  $\rm H_2O$ . The reaction To 3.50 g (13.9 mmol) of 3-aminomethyl-1-benzhydryl ether. The combined organic layer was washed with brine, dried with Na<sub>2</sub>SO<sub>4</sub> and of the title compound.

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(iii) 3-(N-tert-butyloxycarbonylaminomethyl) azetidine 35

3.4 g (9.6 mmol) of 3-(N-tert-

butyloxycarbonylaminomethyl)-1-benzhydryl azetidine was  $\mathrm{Pd}(\mathrm{OH})_2$  at 5 MPa over night. The catalyst was filtered dissolved in 170 mL MeOH and hydrogenated over 0.30 g off and the solvent evaporated. The crude product was

(g))/ $\mathrm{CH}_2\mathrm{Cl}_2$ , 1/9, as eluent to yield 1.2 g (67 %) of the  $\mathrm{MeOH/CH_2Cl_2}$ , 1/9, followed by MeOH (saturated with NH<sub>3</sub> purified by flash chromatography using title compound.

(iv) 3-(N-tert-butyloxycarbonylaminomethyl)-1-(Nbenzyloxycarbonylamidino) azetidine (Boc-Mig( $\mathbb{Z}$ )) 2

mmol) of N-benzyloxycarbonyl-O-methylisourea was mixed in 6.5 mL toluene and heated to 70°C for 72 h and then by MeOH (saturated with NH3(g))/CH2Cl2, 1/9, as eluent left at roomtemperature for another 72 h. Evaporation followed by flash chromatography using EtOAc followed butyloxycarbonylaminomethyl) azetidine and 1.3 g (6.3 gave 0.67 g (38 %) of the title compound as a white 0.9 g (4.8 mmol) of 3-(N-tert-

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(v) 3-aminomethyl-1-(N-benzyloxycarbonylamidino) azetidine (H-Mig(Z))

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0.67 g (1.85 mmol) of Boc-Mig(2) was dissolved in 10 mL dried with  $\mathrm{Na}_2\mathrm{SO}_4$  and evaporated to yield 0.43 g (89 %) of EtOAc saturated with HCl(g) and stirred for 10 min. and the aqueous phase was extracted with  $3x8\,$  mL EtOAc. KOH(aq) was added dropwise. The layers were separated at roomtemperature. 10 mL of a saturated solution of The organic layers were combined, washed with brine, of the title compound.

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2H), 3.66 (dd, 2H) 4.03 (dd, 2H) 5.07 (s, 2H), 7.2-7.4 <sup>1</sup>H-NYR (300 MHz, CDCl<sub>3</sub>): 6 2.55-2.65 (m, 1H), 2.84 (d, 32

MS m/z 263 (M+ + 1)

3-aminoethyl-1-(N-benzyloxycarbonylamidino) azetidine

prepared according to the literature, see A.G. Anderson, Jr., and R. Lok, J.Org.Chem., 37, 3953, 1972. (i) 3-carboxylic acid-1-benzhydryl azetidine was

# (ii) 3-hydroxymethyl-1-benzhydryl azetidine

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as pale yellow crystals. of the solvent gave 7.1 g (86 %) of the title compound filter cake was washed repeatedly with THF. Evaporation  $\mathrm{NH_4Cl}\left(\mathrm{aq}\right)$  , the gelatinous mixture was filtered and the hydrolyzed by careful addition, with cooling, of was refluxed for 3.5 h. Excess hydride reagent was in 30 mL THF at roomtemperature. The reaction mixture slowly to a suspention of 4.9 g (130.2 mmol) of  ${\tt LiAlH}_4$ benzhydryl azetidine in 80 mL of dry THF was added A solution of 8.7 g (32.5 mmol) 3-carboxylic acid-1-

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# (iii) 3-methanesulfonatomethyl-1-benzhydryl azetidine

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30 25 reaction mixture was poured into a mixture of ice and  $m H_2O$ . The precipitate was collected, washed with  $m H_2O$  and allowed to stand in a refrigerator over night. The dried in vacuo to yield 7.75 g (89.5 \$) of the title The reaction mixture was stirred for 1 h. and then 4.50 g (39.2 mmol) of methanesulfonyl chloride at 0°C. benzhydryl azetidine in 50 mL of dry pyridine was added To a solution of 6.62 g (26.1 mmol) 3-hydroxymethyl-1-

# (iv) 3-cyanomethyl-1-benzhydryl azetidine

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To a solution of 7.75 g (23.4 mmol) 3-

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ഗ vacuo to yield 5.7 g (93 %) of the title compound. precipitate was collected, washed with  $\mathrm{H}_2\mathrm{O}$  and dried in cooled, and poured into a mixture of ice and  $\mathrm{H}_2\mathrm{O}$ . The 10 mL  ${
m H_2O}$ . The mixture was heated at 65°C for 20 h, DMF was added a solution of 3.44 g (70.0 mmol) NacN in methanesulfonatomethyl-1-benzhydryl azetidine in 50 mL

## (v) 3-aminoethyl-1-benzhydryl azetidine

- 20 15 10 gave 5.0 g (87 %) of the title compound. brine and dried with  ${
  m Na_2SO_4}.$  Evaporation of the solvent residue was dissolved in diethyl ether, washed with mixture was filtered and the filter cake was washed repeatedly with THF. The solvent was evaporated, the addition, with cooling, of  $\mathrm{NH_4Cl}\left(\mathrm{aq}\right)$ , the gelatinous 4 h. Excess hydride reagent was hydrolyzed by careful 5.7 g (21.7 mmol) of 3-cyanomethyl-1-benzhydryl roomtemperature. The reaction mixture was refluxed for azetidine was added slowly to a suspention of 2.9 g (76.0 mmol) of LiAlH $_4$  in 80 mL of dry THF at
- (vi) 3-(N-tert-butyloxycarbonylaminoethyl)-1-benzhydryl
- 25 azetidine, in a yield of 6.5 g (95 %). tert-butyloxycarbonylaminomethyl)-1-benzhydryl benzhydryl azetidine according the procedure for 3-(N-The title compound was prepared from 3-aminoethyl-1-
- 30 (vii) 3-(N-tert-butyloxycarbonylaminoethyl) azetidine

according the procedure for 3-(N-tertbutyloxycarbonylaminoethyl)-1-benzhydryl azetidine The title compound was prepared from 3-(N-tert-

butyloxycarbonylaminomethyl) azetidine, in a yield of 1.2 9 (70 %).

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(viii) 3-(N-tert-butyloxycarbonylaminoethyl)-1-(Nbenzyloxycarbony lamidino) azetidine (Boc-Dig(Z)) The title compound was prepared from 3-(N-tert-butyloxycarbonylaminoethyl) azetidine according the procedure for 3-(N-tert-butyloxycarbonylaminomethyl)-1-(N-benzyloxycarbon ylamidino) azetidine, in a yield of 0.090 g (34 %).

10 (ix) 3-aminoethyl-1-(N-benzyloxycarbonylamidino)
azetidine (H-big(2))

0.589 g (1.56 mmol) of Boc-Dig(2) was dissolved in 10 mL of EtoAc saturated with HCl(g) and stirred for 10 min. at roomtemperature. 10 mL of a saturated solution of KOH(ag) was added dropwise. The layers were separated and the aqueous phase was extracted with 3x8 separated and the aqueous phase was extracted with at EtoAc. The organic layers were combined, washed with brine, dried with Na<sub>2</sub>SO<sub>4</sub> and evaporated to yield 0.415 g (96 %) of the title compound.

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<sup>1</sup>H-NYR (500 MHz, CDCl<sub>3</sub>): 6 1.60 (dt, 2H), 2.52-2.54 (m, 3H), 3.53 (bs, 2H), 4.0 (bt, 2H), 5.00 (s, 2H), 7.17-7.31 (m, 5H).

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Working Examples

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Example 1

HOOC-CH2-(R) Cg1-Aze-Pab

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(1) Boc-(R)Cgl-Aze-Pab(Z)

To a stirred mixture of 3.40 g (10 mmol) Boc-(R)Cgl-Aze-OH (See Preparation of starting materials) and 5.13 g DNAP (42 mmol) in 120 ml CH<sub>3</sub>CN was added 3.18 g H-Pab(Z) x HCl (See Preparation of starting materials).

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title compound.

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After stirring for 2 hours at room temperature the mixture was cooled to -8°C and 2.01 g (10.5 mmol) EDC was added. The reaction was allowed to reach room temperature and the stirring was continued for an additional 47 hours. The solvent was evaporated and the residue was dissolved in 200 ml EtOAc. The organic phase was washed with 1 x 50 ml water, 1 x 50 + 2 x 25 ml 0.5 M KHSO<sub>4</sub>, 2 x 25 ml NaHCO<sub>3</sub>(saturated), 1 x 50 ml water and dried. Evaporation of the solvent gave 5.21 g water and dried. Evaporation of the solvent gave 5.21 g

1H-NWR (500 MHz, CDCl<sub>3</sub>): 6 0.8-1.9 (m, 20H; thereof 1.30 (s, 9H)), 2.35-2.6 (m, 2H), 3.74 (bt, 1H), 4.10 (m, 1H), 4.25-4.4 (m, 2H), 4.45-4.6 (m, 1H, rotamers), 4.75-5.0 (m, 1H, rotamers), 5.08 (bd, 2H), 5.15 (s, 2H), 7.15-7.35 (m, 5H), 7.41 (d, 2H), 7.77 (d, 2H), 8.21 (m, 1H).

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(11) H-(R)Cgl-Aze-Pab(Z)

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To a cold (ice bath temperature) solution of 18.8 g HCl<sub>(g)</sub> in 195 ml EtOAc was added 4.69 g (7.743 mmol) of Boc-(R)Cgl-Aze-Pab(Z) together with 40 ml EtOAc. The reaction mixture was allowed to reach room temperature and stirred for 30 min. 140 ml Et<sub>2</sub>O was added to the clear solution where upon a precipitate was formed. The reaction was left at room temperature for an additional 1 h and 40 minutes. The precipitate was filtered off, washed quickly with 150 ml Et<sub>2</sub>O and dried in vaccuo. The precipitate was dissolved in 50 ml of water and made alkaline with 15 ml 2 M NaOH. The alkaline waterphase was extracted with 1'x 100 + 1 x 50 ml cH<sub>2</sub>Cl<sub>2</sub>. The combined organic phase was washed with 1 x 20 ml water, 1 x 20 ml Brine and dried(MgSO<sub>4</sub>). Evaporation of the solvent gave 3.44 g (88%) of the

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1H), 4.43 (dd, 1H), 4.53 (dd, 1H), 4.91 (m, 1H), 5.22 1H), 2.67 (m, 1H), 3.07 (d, 1H), 4.11 (m, 1H), 4.18 (m, (s, 2H), 7.2-7.4 (m, 7H), 7.45 (d, 2H), 8.51 (d, 2H).  $^{1}\text{H-NMR}$  (500 MHz, CDCl<sub>3</sub>): 6 0.8-2.0 (m, 11H), 2.51 (m,

(iii) BnOOC-CH<sub>2</sub>-(R)Cgl-Aze-Pab(Z)

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eluents to give 0.525 g (36 %) of the title compound. 95/5 and then diethylether/MeOH(NH $_3$ -saturated) 9/1 as chromatography using first  $CH_2Cl_2/MeoH(NH_3-saturated)$ g of a residue that was twice subjected to flash EtOAc. The organic phase was washed with water, dried  $(\mathrm{Na_2SO_4})$  , filtered and evaporated in vacuo to give 1.17 alcaline with 1 N NaOH to pH>8 and extracted with was extracted with 1 M  $ext{KHSO}_4$  and this waterphase was washed with EtOAc. The acidic waterphase was made the mixture was washed with water, the organic phase solvent was evaporated in vacuo. EtOAc was added and were mixed and heated in a 60°C oilbath for 3 h. The materials), 0.99 g (5.6 mmol)  $\mathrm{K}_2\mathrm{CO}_3$  and 113 ml  $\mathrm{CH}_3\mathrm{CN}$  $\mathrm{NO_2})\,\mathrm{Ph}\mathrm{-SO_2}\mathrm{-OCH_2}\mathrm{-COOBn})$  (See Preparation of starting benzyl-2-(orto-nitrobenzenesulfonyloxy)acetate ((2-1.13 g (2.2 mmol) H-(R)Cgl-Aze-Pab(Z), 0.9 g (2.6 mmol)

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using the same procedure as above to give the title  ${\rm OCH_2 extsf{-}COOBn})$  (See Preparation of starting materials) compound in 52 % yield. The alkylation was also carried out using Benzyl-2-(para-nitrobenzenesulfonyloxy)acetate ((4-NO<sub>2</sub>)Ph-SO<sub>2</sub>-

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(m, 10H), 7.45 (d, 2H), 7.79 (d, 2H), 8.42 (m, 1H). 1H), 4.91 (m, 1H), 5.07 (s, 2H), 5.22 (s, 2H), 7.2-7.4 1H), 3.95 (m, 1H), 4.05 (m, 1H), 4.44 (m, 1H), 4.55 (m, 1H), 2.63 (m, 1H), 2.88 (d, 1H), 3.24 (d, 1H), 3.27 (d, <sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>): & 0.85-2.15 (m, 11H), 2.48 (m,

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(iva)  $HOOC-CH_2-(R)Cgl-Aze-Pab \times 2 HCl$ 

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The state of

ഗ chloroform and 5 \$ Pd/C were added and the mixture was from water to give 11 mg (72%) of the title compound. filtration and evaporation the product was lyophilized hydrogenated at atmospheric pressure for 1 h. After dissolved in 5 ml of methanol. A few drops of  $Bnooc-cH_2-(R)cgl-Aze-Pab(Z)$ , 20 mg (0.031 mmol), was

10 4.4-4.55 (m, 2H), 4.66 (s, 2H), 5.08 (m, 1H), 7.65 (d, 2.44 (m, 1H), 2.82 (m, 1H), 3.90 (s, 2H), 4.09 (d, 1H), 2H), 7.89 (d, 2H). <sup>1</sup>H-NMR (500 MHz, D<sub>2</sub>O): & 1.0-2.0 (m, 11H), 2.10 (m, 1H),

8 167.3, 167.9, 169.9 and 172.4.  $^{13}\mathrm{C\text{-}NMR}$  (75.5 MHz, D<sub>2</sub>0): amidine and carbonyl carbons:

(ivb) HOOC-CH<sub>2</sub>-(R)Cgl-Aze-Pab

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20 for 5 hours. Filtration of the catalyst through cellite and evaporation of the solvent gave the title compound in and hydrogenated over 5 % Pd/C at atmospheric pressure BnOOC-CH2-(R)Cgl-Aze-Pab(Z) was dissolved in EtoH (99%)

30 25 7.54 (d,2H), 7.66-7.71 (d, 2H). 1H), 4.43-4.60 (AB-system, 2H), 4.80-4.85 (dd, 1H), 7.48system plus d, 3H), 4.18-4.25 (bq, 1H), 4.28-4.32 (bq, 2.32-2.42 (m, 1H), 2.54-2.64 (m, 1H), 2.95-3.10 1.70 (m, 3H), 1.73-1.85 (m, 2H), 1.94-2.02 (bd, 1H), rotamer: 6 1.00-1.12 (m, 1H), 1.13-1.34 (m, 4H), 1.55- $^{
m 1}H ext{-NMR}$  (500 MHz, CD $_{
m 3}$ OD, mixture of two rotamers): major (AB-

7.57 (bd), 7.78 (bd). Resolved signals from the minor rotamer appears at  $\delta$  0.95 (m), 1.43 (m), 2.24 (m), 2.84 (d), 3.96 (m), 4.03 (m),

35  $^{13}\mathrm{C\text{-}NMR}$  (125 MHz, CD $_3$ OD): amidine and carbonyl carbons:  $\delta$ 168.0, 173.0, 176.3 and 179.0

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#### Example 2

HOOC-CH2-CH2-(R) Cgl-Ase-Pab x 2 HCl

(i) H-(R)Cgl-Aze-Pab(Z)

prepared in the same way as decribed in Example 1 (11) by treating the formed hydrochloride salt with base to afford the free base.

(ii) Bnooc-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cgl-Aze-Pab(Z)

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H-(R)Cgl-Aze-Pab(2), 0.19 g (0.38 mmol), and 70 mg ( 0.43 isopropanol. The mixture was left standing for 6 days. Flash chromatography using  $\mathrm{CH_2Cl_2/THF} = 8/2$  as eluent mmol) of benzyl acrylate were dissolved in 2 ml of afforded 0.12 g (48%) of the title compound.

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1H NWR (500 MHz, CDC13) & 0.8-1.9 (m, 10 H), 1.95 (bd, 1 H), 2.4-2.6 (m, 4 H), 2.7-2.8 (m, 3 H; thereof 2.79 (d, 1 H)), 4.13 (m, 1 H), 4.37 (dd, 1 H), 4.60 (dd, 1 H), 4.97 (dd, 1 H), 5.09 (dd, 2 H), 5.22 (s, 2 H), 7.25-7.4 (m, 10 H), 7.47 (d, 2 H), 7.83 (d, 2 H), 8.61 (bt, 1 H). 20

(iii)  $HOOC-CH_2-CH_2-(R)Cgl-Aze-Pab \times 2 HCl$ 25

BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cgl-Aze-Pab(Z), 0.10 g (0.15 mmol), was dissolved in 10 ml of ethanol and hydrogenated over 5% Pd/C at atmospheric pressure for 1 h.

product was purified on RPLC using  $\mathrm{CH_3CN/0.1}$  M  $\mathrm{NH_4OAc}$ with an excess of conc. Hcl and freeze dried again to The solution was filtered, evaporated and the crude (1/4). The resulting product was freeze dried, treated give 31 mg of the dihydrochloride salt.

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2.7-2.9 (m, 3 H), 3.2-3.4 (m, 2 H), 3.98 (d, 1 H), 4.35- $^{1}\mathrm{H}$  NMR (300 MHz,  $^{1}\mathrm{D_{2}O})$  6 0.8-2.1 (m, 11 H), 2.38 (m, 1 H),

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4.55 (m, 2 H), 4.60 (s, 2 H), 5.04 ( dd, 1 H), 7.59 (d,

2 H), 7.83 (d, 2 H).

 $^{13}\mathrm{C}$  NMR (75 MHz,  $\mathrm{D_2O}$ ): amidine and carbonyl carbons:  $\delta$ 167.2, 167.8, 172.3 and 175.5.

#### Example 3

HOOC-CH2-(R) Cgl-Pro-Pab x 2 HCl

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(i) Boc-(R)Cgl-Pro-Pab(Z)

acetate. The organic phase was washed twice with a 0.3 M stirred over night. After evaporation , the residue was acidic water phase was extracted three times with ethyl  $\mathrm{KHSO_4}\mathrm{-solution}$ , twice with a  $\mathrm{NaHCO_3}\mathrm{-solution}$  and once with brine, dried (Na2SO4), filtered and evaporated. The crude product was purified by flash chromatography on silica gel using ethyl acetate as eluent to yield 1.77 g ( 44% 1.31 g (6.81 mmole) was added. The temperature was allowed to reach room temperature and the mixture was dissolved in ethyl acetate and 0.3 M  $\rm KHSO_4$ -solution. The starting materials), 2.3 g (6.49 mmol) , DMAP, 2.38 g (19.47 materials), 1.84 g (6.49 mmol) was mixed in 30 ml acetonitrile. The mixture was cooled to -15°C and EDC, and H-Pab(Z)(See preparation of ō preparation (See Boc-(R)Cgl-Pro-OH ) of the product. mmol),

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2H), 7.18-7.4 (m, 5H), 7.45 (d, 2H), 7.62 (bs, 1H), 7.81 9H), 2.37 (bs, 1H), 3.53 (g, 1H), 3.94 (bs, 1H), 4.02 (m, 1H), 4.43 (bs, 2H), 4.65 (d, 1H), 5.09 (bs, 1H), 5.20 (s, H-NMR (500 MHz, CDCl3): 6 0.9-1.49 (m, 14H), 1.5-2.1 (m, (m, 2H),

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(ii) H-(R)Cgl-Pro-Pab(Z)

dihydrochloride salt of the product. solvent was evaporated to yield 1.3 allowed to stand for 10 min at room temperature. The in 75 ml HCl saturated ethyl acetate. The mixture was 1.45 g (2.34 mmol) of Boc-(R)Cgl-Pro-Pab(Z) was dissolved g of

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4.5-4.66 (m, 3H), 5.49 (s, 2H), 7.45-7.7 (m, 7H), 7.87 9H), 2.3-2.5 (m, 1H), 3.75-3.90 (m, 2H), 4.25 (d, 2H), <sup>1</sup>H-NMR (300 MHz, D<sub>2</sub>0): 6 1.0-1.45 (m, 5H), 1.58-2.2 (m,

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evaporated to yield 1.19 g (97%) of the title compound. was washed once with brine, dried ( ${
m Na}_2{
m SO}_4$ ), filtered and phase three times with ethyl acetate. The organic phase salt in 0.1 M NaOH-solution and extracting the water The amine was obtained by dissolving the dihydrochloride

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## (iii) BnOOC-CH<sub>2</sub>-(R)Cgl-Pro-Pab(Z)

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from the column followed by 133 mg (31%) of the desired product Bn00C-CH<sub>2</sub>-(R)Cgl-Pro-Pab(Z). (9%) of  $(BnOOC-CH_2)_2-(R)Cgl-Pro-Pab(2)$  which eluated first acetate/toluene (9/1, 93/7, 95/5, 100/0) to give 46 mg chromatography using a stepwise gradient of ethyl TIC. The mixture was therefore purified further by flash to give 299 mg of a mixture of two products according to stepwise gradient of  $CH_2Cl_2/MeOH$  (97/3 followed by 95/5) product was purified by flash chromatography using a brine, dried  $(\mathrm{Na_2SO_4})$ , filtered and evaporated. The crude and the organic layer was washed once with water and temperature. The reaction mixture was diluted with  $\mathrm{CH_2Cl_2}$ reaction mixture was then stirred over night at room dichloromethane and refluxed for half an hour. The starting materials), 0.299 g ( 2.17 mmole )  ${
m K_2CO_3}$  in 4 ml 0.215 g ( 0.65 mmole ) BnOOC-CH $_2$ -OTf (see preparation of 0.340 g ( 0.65 mmole ) H-(R)Cgl-Pro-Pab(2) was mixed with

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5.19 (s, 2H), 6.75 (bs, NH), 7.1-7.5 (m, 12H), 8.7-8.8 4.43, 2 H), 4.62 (d, 1H), 4.91 (apparent singlet, 2H), 3.5-3.6 (m, 2H), 4.29-4.57 (ABX-system centered at d (m, 2H+NH), 9.45 (bs, NH) 0.9-1.3 (m, 5H), 1.4-2.1 (m, 9H), 2.3-2.4 (m, 1H), 3.05  $^{1}\text{H-NMR}$  (300 MHz, CDCl<sub>3</sub>): BnOOC-CH<sub>2</sub>-(R)Cg1-Pro-Pab(Z):  $\delta$ (d, 1H), 3.20-3.37(AB-system centered at & 3.29, 2 H),

10 4 H), 5.19 (5, 2H), 6.66 (bs, NH), 7.1-7.5 (m, 17H), 7.75 d3.67, 4 H), 4.38-4.58 (ABX-system centered at d4.48, 2 (d, 2H), 7.80 (t, NH), 9.37 (bs, NH) H), 4.68 (d, 1H), 4.82-4.98 (AB-system centered at d 4.91, 2.0 (m, 7H), 2.05 (bd, 1H), 2.3-2.4 (m, 1H), 3.15 (d, 1H), 3.25-3.48 (m, 2H), 3.55-3.79 (AB-system centered at 6 0.68-0.9 (m, 2H), 1.0-1.3 (m, 3H), 1.43 (bd, 1H), 1.55-1H-NMR (300 MHz, CDCl<sub>3</sub>): (BnOOC-CH<sub>2</sub>)<sub>2</sub>-(R)Cgl-Pro-Pab(Z):

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164.7, 168.1, 171.5, 172.3 and 172.6  $^{13}\mathrm{C-NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $\delta$ 

### 20 (iv) HOOC-CH<sub>2</sub>-(R)Cgl-Pro-Pab x 2 HCl

25 for one hour. After filtration through hyflo and ml ethanol. The mixture was treated under  ${
m H_2 ext{-}atmosphere}$ mg, was obtained by freeze drying twice from water. evaporation of the solvent the product in 90% yield, 93 mixed with 0.060 g 5 % Pd/C, 1ml 1M HCl-solution and 10 0.133 g (0.20 mmole ) of BnOOC-CH $_2$ -(R)Cgl-Pro-Pab(Z) was

30 9H), 2.2-2.4 (m, 1H), 3.55-3.85 (m, 4H; thereof 3.79 (s, 2H)), 4.23 (d, 1H), 4.33-4.57 (m, 3H), 7.44 (d, 2H), 7.69  $^{1}\text{H-NMR}$  (300 MHz,  $D_{2}\text{O}$ ): & 1.0-1.45 (m, 5H), 1.5-2.1 (m,

35 166.9, 167.2, 169.1, 174.5  $^{13}\mathrm{C\text{--}NMR}$  (75 MHz,  $\mathrm{D}_2\mathrm{O}$ ): amidine and carbonyl carbons: §

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HOOC-CH2-CH2-(R) CG1-Pro-Pab x 2 HCl

(1) BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cgl-Pro-Pab(Z)

0.406 g (0.782 mmole) of H-(R)cgl-Pro-Pab(Z) (See Example 3) was dissolved in 3 ml ethanol and 132  $\mu$ l (0.861 mmole) of bensylacrylate was added. The reaction mixture was stirred for three days at room temperature. The mixture was evaporated and the crude product purified by flash chromatography using a stepwise gradient of CH<sub>2</sub>Cl<sub>2</sub>:MeOH 95/5 and 90/10 as eluent to yield 0.399 g (75%) of the product.

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JH-NMR (300 MHz, CDCl<sub>3</sub>):6 0.8-1.0 (m, 1H), 1.0-1.3 (m, 4H), 1.35-2.2 (m, 9H), 2.3-2.6 (m, 4H), 2.65-2.78 (m, 1H), 3.05 (d, 1H), 3.4-3.6 (m, 2H), 4.25-4.52 (ABX-system 1H), 3.05 (d, 1H), 3.4-3.6 (m, 2H), 4.64 (dd, 1H), 5.05 (s, 2H), 5.20 (s, 2H), 7.2-7.38 (m, 10H), 7.43 (d, 2H), 7.78 (d, 2H), 7.2-7.38 (m, 10H), 7.43 (d, 2H), 7.78 (d, 2H), 7.8 (d, 2H), 7

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13C-NMR (75 MHz, CDCl<sub>3</sub>): amidine and carbonyl carbons: 6 164.7, 167.9, 171.3, 172.7 and 175.4.

(ii) HOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cgl-Pro-Pab x 2 HCl

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0.261 g (0.383 mmole) of BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)cgl-Pro-Pab(Z) was mixed with 0.075 g 5 % Pd/C, 1ml 1M HCl-solution and 10 ml ethanol. The mixture was hydrogenated at atmospheric pressure for two hours. After filtration through hyflo and evaporation of the solvent the product through hyflo and evaporation of the solvent the product o.196 g (96%) was obtained by freeze drying twice from

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<sup>1</sup>H-NWR (300 MHz, D<sub>2</sub>O): 6 1.17-1.40 (m, 5H), 1.60-1.92 (m, 5H), 1.92-2.2 (m, 4H), 2.32-2.48 (m, 1H), 2.81 (t, 2H),

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3.11-3.36 (ABX<sub>2</sub>-system centered at 6 3.24, 2H), 3.63-3.90 (m, 2H), 4.25 (d, 1H), 4.42-4.63 (m, 3H), 7.54 (d, 2H), 7.78 (d, 2H)

5 13C-NMR (75 MHz, D<sub>2</sub>0): amidine and carbonyl carbons: 6 167.0, 167.30, 174.6 and 174.7.

Example 5

10 (HOOC-CH2)2-(R)Cg1-Pro-Pab x 2 HCl

46 mg (0.056 mmole) of (Bnooc-CH<sub>2</sub>)<sub>2</sub>-(R)Cgl-Pro-Pab(Z) (See Example 3) was mixed with 25 mg 5 % Pd/C , 0.7 ml 1M HCl-solution and 7 ml ethanol. The mixture was hydrogenated at atmospheric pressure for one hour. The catalyst was filtered off through hyflo and the solvent evaporated. The final product 25 mg (77 %) was obtained by freeze drying twice from water.

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20 <sup>1</sup>H-NWR (300 MHz, D<sub>2</sub>O): 6 1.0-1.4 (m, 5H), 1.45-2.2 (m, 9H), 2.25-2.45 (m, 1H), 3.53-3.84 (m, 2H), 3.84-4.22 (AB-system centered at 6 4.03, 4 H), 4.26 (d, 1H), 4.35-4.6 (m, 3H), 7.53(d, 2H), 7.77 (d, 2H)

 $^{13}\text{C-NMR}$  (75 MHz,  $D_2\text{O})\colon$  amidine and carbonyl carbons: 6 167.1, 167.3, 170.6 and 174.5

Example 6

30 H-(R)Cgl-Pic-Pab x 2 HCl

(i) Boc-(R)Cgl-Pic-Pab(Z)

0.478 g (2.49 mmol) EDC was added at -18°C to a stirred solution of 0.875 g (2.37 mmol) Boc-(R)Cgl-Pic-OH (See preparation of starting materials), 1.22 g (9.97 mmol) DMAP, and 0.706 g (2.49 mmol) H-Pab(Z) (See preparation

of starting materials) in a mixture of 30 ml acetonitrile and 1 ml DMF. The reaction mixture was allowed to reach room temperature during a couple of hours and stirring was continued for 48 h. The solvent was removed in vacuo and the residue was dissolved in 50 ml ethyl acetate. The solution was washed with 15 ml water, 3x15 ml 0.3 M KHSO<sub>4</sub>, 2x15 ml Na<sub>2</sub>CO<sub>3</sub> solution and water. Removal of the solvent gave a residue which was subjected to flash chromatography using ethyl acetate/heptane 9:1 as eluent. The yield was 0.96 g (64%).

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## (ii) H-(R)Cgl-Pic-Pab(Z)

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Hydrogen chloride was bubbled through a solution of 0.56 g (0.88 mmol) Boc-(R)Cgl-Pic-Pab(Z) in 25 ml ethyl acetate. After a couple of minutes crystals precipitated from the solution. The solvent was removed in vacuo and 50 ml ethyl acetate was added. Washing with 2x15 ml 2 M sodium hydroxide solution and extraction of the aqueous phase with 25 ml ethyl acetate was followed by drying (sodium sulphate) of the combined extracts and removal of the solvent in vacuo to give 0.448 g (95%) of the desired product.

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## 25 (iii) H-(R)Cgl-Pic-Pab x 2 HCl

A solution of 98 mg (0.18 mmol) H-Cgl-Pic-Pab(Z) in 5 ml 95% ethanol and 1 ml water was stirred in an atmosphere of hydrogen for 4 hours in the presence of 5% Pd/C. The mixture was filtered and 0.3 ml 1 M hydrochloric acid was added. The ethanol was removed in vacuo and the residue was freeze dried to give 70 mg (81%) of the desired compound.

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2H), 7.76 (d, 2H)

 $^{13}\text{C-MMR}$  (75 MHz,  $D_20\}$ : amidine and carbonyl carbons:  $\emph{b}$  167.2, 170.5 and 173.4.

#### Example 7

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HOOC-CH2-(R,8)CH(COOH)-(R)CG1-Pic-Pab x 2 HC1

(i) Bnooc-CH<sub>2</sub>-(R,S)CH(COOBn)-(R)Cgl-Pic-Pab(Z)

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A mixture of 350 mg (0.66 mmol) H-(R)Cgl-Pic-Pab(Z) (See Example 6) and 233 mg dibenzyl maleate in 2.5 ml ethanol was kept at room temperature for 4 days. The ethanol was removed in vacuo and the residue was subjected to flash chromatography using ethyl acetate/heptane 9:1 as eluent to give 0.108 mg of the product.

# (ii) $HOOC-CH_2-(R,S)CH(COOH)-(R)Cgl-Pic-Pab \times 2 HCl$

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105 mg (0.13 mmol) Bn00C-CH<sub>2</sub>-(R,S)CH(COOBn)-(R)Cgl-Pic-Pab(Z) dissolved in 5 ml 95% ethanol and 1 ml water was hydrogenated for 5 hours in the presence of 5 % Pd/C. 0.3 ml 1 M hydrochloric acid was added and the mixture was filtered and the solvent was removed in vacuo. The residue was dissolved in water and freeze dryed to yield 54 mg (73%) of the desired substance.

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 $^{13}\text{C-NMR}$  (75 MHz  $D_2\text{O}$ ): amidine and carbonyl carbons:  $\delta$  167.1, 168.95, 169.6 and 173.1.

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MS m/z 516 (M+ +1)

Example 8

H-(R) Cha-Ase-Pab x 2 HCl

(i) Boc-(R)Cha-Aze-Pab(Z)

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and the solvent was subsequently removed in vacuo. The 3x10 ml 0.3M KHSO4, 2x10 ml  ${\rm Na_2CO_3\text{-}NaCl}$  (aq), and finally 10 ml Brine. Drying ( $Na_2SO_4$ ) and removal of the solvent in vacuo gave a residue which was subjected to flash chromatography using ethyl acetate/methanol 9:1 as eluent starting materials) in 20 ml acetonitrile. The reaction mixture was allowed to reach room temperature over night residue was dissolved in 40 ml ethyl acetate and the mixture of 0.72 g (2.03 mmol) Boc-(R)Cha-Aze-OH (See DMAP, and 604 mg (2.13 mmol) H-Pab(2) (See preparation of organic phase was washed succesively with 10 ml water, 409 mg (2.13 mmol) EDC was added at -18°C to a stirred preparation of starting materials), 1.04 g (8.53 mmol) to yield 645 mg (51%) of the title compound.

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(ii) H-(R)Cha-Aze-Pab(Z)

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dried (Na<sub>2</sub>CO<sub>3</sub>) and the solvent was removed in vacuo to applied to remove excess hydrogen chloride and the Washing with 2x15 ml Na<sub>2</sub>CO<sub>3</sub> (aq) was followed by The combined organic extracts were washed with water and acetate. After a couple of minutes, TLC analysis indicated the completion of the reaction. Vacuum was mixture was then diluted to 50 ml with ethyl acteate. extraction of the aqueous phase with 15 ml ethyl acetate. Hydrogen chloride was bubbled through a solution of 640 mg (1.03 mmol) Boc-(R)Cha-Aze-Pab(Z) in 25 ml of ethyl give 513 mg (96%) of  $H^{-}(R)$ Cha-Aze-Pab(Z).

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(iii) H-(R)Cha-Aze-Pab x 2 HCl

76 mg (0.15 mmol) H-(R)Cha-Aze-Pab(Z) dissolved in 5

IM hydrochloric acid and evaporation of the solvent in 95% ethanol and 1 ml water was hydrogenated at atmospheric pressure in the presence of 5% Pd/C for 4 h. Removal of the catalyst by filtration, addition of 0.4 ml vacuo gave a residue which was dissolved in 2 ml water. Freeze drying gave 57 mg (85%) of the product. <sup>1</sup>H-NMR (500 MHz, D<sub>2</sub>O, 2 rotamers, 3:1 mixture): 6 1.02-2.80-2.90 (m, 1H), 4.25 (bt, 1H), 4.40 (dg, 1H), 4.53 1.20 (m, 2H), 1.22-1.92 (m, 11H), 2.40-2.50 (m, 1H), (dq, 1H), 4.65 (s, 2H), 5.05-5.10 (m, 1H), 7.65 (d, 2H),

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7.88 (d, 2H).

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Chemical shifts of resolved signals of the minor rotamer: 8 0.57 (m), 0.85 (m), 2.95 (m), 4.06 (dq), 4.17 (dq), 4.63 (s), 5.33(m), 7.70(d), 7.93 (d).  $^{12}\text{C-NMR}$  (125 MHz  $^{} ext{D}_2 ext{O} ext{)}:$  amidine and carbonyl carbons:  $^{\delta}$ 167.2, 170.4 and 172.8.

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Example 9

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HOOC-CH2-(R) Cha-Aze-Pab x 2 BC1

(i) BnOOC-CH<sub>2</sub>-(R)Cha-Aze-Pab(Z)

was washed with brine and dried  $(\mathrm{Na_2SO_4})$  . Evaporation in 0.119 g (0.52 mmol) benzyl bromoacetate was added to a mixture of 0.27 g (0.52 mmol) H-(R)Cha-Aze-Pab(Z) (See Example 8) and 0.158 g. (1.14 mmol)  $\rm K_2CO_3$  in 5.2 ml acetonitrile and heated to 60°C in an oilbath for 1 h. The solvent was removed and ethyl acetate and water was added. The phases were separated and the organic phase 35 30

vacuo gave 0.344 g of a residue which was subjected to

saturated methanol (60:5:2) to give 0.163 g of the then another time using ethylacetate:tetrahydrofuran:  $\mathrm{NH_3}$ desired product. flash chromatography using ethyl acetate as eluent, and

12H), 7.7-7.85 (d, 2H), 8.3-8.45 (t, 1H). 4.8-4.95 (m, 1H), 5.05 (s, 2H), 5.2 (s, 2H), 7.2-7.5 (m, 3H), 3.95-4.05 (t, 2H), 4.4 and 4.5 (ABX-system, 2H), 11H), 2.35-2.55 (m, 1H), 2.55-2.75 (m, 1H), 3.15-3.32 (m, 1H-NMR (300 MHz, CDCl<sub>3</sub>); & 0.7-1.0 (m, 2H). 1.05-2.05 (m,

164.5, 167.8, 170.7, 171.9 and 175.9.  $^{13}\text{C-NMR}$  (75 MHz, CDCl<sub>3</sub>): amidine and carbonyl carbons: 6

## (ii) HOOC-CH<sub>2</sub>-(R)Cha-Aze-Pab x 2 HC]

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water and freeze drying gave 107 mg (85 %) of the title evaporation of the solvent followed by dissolving in Pd/C for 4 h. Removal of the catalyst by filtration and in 5.5 ml ethanol (99.5 %) and 0.7 ml hydrogen chloride (1 N) was hydrogenated in the presence of 0.17 g 5 \$0.163 g (0.243 mmol)  $BnOOC-CH_2-(R)Cha-Aze-Pab(Z)$  dissolved

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25 Resolved signals from the minor rotamer appears at  $\delta$ (m, 2H). (m, 1H), 4.83-4.88 (m, 1H), 7.5-7.6 (m, 2H), 7.73-7.82 (m, 1H), 4.36-4.43 (m, 1H), 4.43-4.5 (m, 1H), 4.58-4.65 (m, 1H), 3.5-3.75 (m, 2H), 4.05-4.15 (m, 1H), 4.15-4.23 rotamer: & 0.95-1.95 (m, 13H), 2.3-2.4 (m, 1H), 2.6-2.75  $^{
m 1}H ext{-NMR}$  (500 MHz, CD $_{
m 3}$ OD, mixture of two rotamers): major

2.2-2.3 (m), 3.95-4.05 (m), 5.1-5.17 (m), 7.6-7.67 (m).

168.2, 169.8, 168.9 and 172.3.  $^{13}\mathrm{C-NMR}$  (75 MHz, CD $_3$ OD): amidine and carbonyl carbons; 6

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#### Example 10

# HOOC-CH2-(R,8)CH(COOH)-(R)Cha-Aze-Pab x 2 HCl

# (i) Bn00C-CH<sub>2</sub>-(R,S)CH(C00Bn)-(R)Cha-Aze-Pab(Z)

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acetate/methanol 95/5 as eluent to give 54 mg (15%) of residue was subjected to flash chomatography using ethyl 5 days. After removal of the ethanol in vacuo, the 1.5 ml 95% ethanol was stirred at ambient temperature for the product. Example 8) and 144 mg (0.487 mmol) dibenzyl maleate in A mixture of 230 mg (0.443 mmol) H-(R)Cha-Aze-Pab(Z) (See

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# (ii) HOOC-CH<sub>2</sub>-(R,S)CH(COOH)-(R)Cha-Aze-Pab

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Removal of the catalyst by filtration and evaporation of drying gave 32 mg (93%) of the product. in 2 ml water and 0.2 ml 1M hydrochloric acid. Freeze the solvent in vacuo gave a residue which was dissolved hydrogenated in the presence of 5% Pd/C for 4.5 h. Pab(2) dissolved in 5 ml 95% ethanol and 1 ml water was 49 mg (0.06 mmol) Bn00C-CH<sub>2</sub>-(R,S)CH(COOBn)-(R)Cha-Aze-

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appears in the spectrum. minor rotamer, integrating to approximatly 15% also two diastereomers. Additionally resolved resonances of a two sets of strongly overlapping signals arising from the The  $^1\mathrm{H-NMR}$  spectrum of the title compound in D $_2\mathrm{O}$  exhibits

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2H), 7.80-7.94 (m, 2H) <sup>1</sup>H-NWR (300 MHz, D<sub>2</sub>0): & 1.03-2.00 (m, 13H), 2.32-2.53 (m, 4H), 4.62 (bs, 2H), 5.00-5.10 (m, 1H), 7.55-7.68 (m, (m, 1H), 2.72-2.96 (m, 1H), 3.06-3.28 (m, 2H), 4.10-4.55

35 Resolved signals from the minor rotamer appears at  $\delta$  0.65 (m), 0.80 (m), 4.00 (m), 5.24 (m), 5.35 (m).

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13C-NMR (75 MHz D2O): amidine and carbonyl carbons: \$ 167.2, 169.0, 171.0, 172.3 and 174.1.

#### Example 11

# HOOC-CH2-(Rors) CH(COOH)-Cha-Are-Pab/a x 2 HCl

# (1) BnOOC-CH<sub>2</sub>-(RorS)CH(COOBn)-(R)Cha-Aze-Pab(Z)/a

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water dried ( ${
m Na}_2{
m SO}_4$ ), filtered and evaporated to yield acetate/methanol 98/2 as eluent to give 1.024 g (32%) of diastereomers were separated by RPLC using (CH3CN/0.1 M  $\mathsf{NH}_{\mathsf{q}}\mathsf{OAC}$  65/35) as eluent. This diastereomer eluted first from the column. After removal of the acetonitrile in vacuo the water phase was extracted three times with ethyl acetate. The organic phase was washed once with After removal of the ethanol in vacuo, the residue was A mixture of 2.0 g (3.8491 mmol) H-(R)Cha-Aze-Pab(Z)(See Example 8) and 1.37 g dibenzyl maleate in 10 ml 95% ethanol was stirred at ambient temperature for 4 days. 0.352 g of the title compound as a pure stereoisomer. using Bnooc-CH2-(R, S) CH(COOBN)-(R) Cha-Aze-Pab(Z). subjected to flash chromatography 20

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# (ii) HOOC-CH2-(RorS) CH(COOH)-(R) Cha-Aze-Pab/a 2 x HCl

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a residue which was dissolved in 5 ml water and 1.0 ml 1M presence of 5% Pd/C for 4.5 h. Removal of the catalyst by filtration and evaporation of the solvent in vacuo gave hydrochloric acid. Freeze drying gave 214 mg (87%) of the  $\operatorname{Pab}(\mathbf{Z})/\mathbf{a}$  (The diaststereomer from (1) above) dissolved in 15 ml 95% ethanol and 3 ml water was hydrogenated in the 350 mg (0.43 mmol) BnOOC-CH<sub>2</sub>-(RorS)CH(COOBn)-(R)Cha-Azeproduct as a pure stereoisomer.

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1H), 4.56 (AB-system, 2H), 4.76-4.86 (m, 1H, partially 1.93 (m, 13H), 2.25-2.38 (m, 1H), 2.60-2.75 (m, 1H), 2.88 (dd, 2H), 3.92 (t, 1H), 4.15-4.25 (m, 2H), 4.30-4.43 (m, obscured by the solvent signal), 7.59 (d, 2H), 7.78 (d, H-NMR (300 MHz, MeOD, mixture of two rotamers): 6 0.85-2H). Resolved signals arising from the minor rotamer appears at 6 0.70, 2.95, 3.82, 4.00, 5.08 and 7.83  $^{13}\mathrm{C-NMR}$  (75 MHz  $\mathrm{D_2O}$ ): amidine and carbonyl carbons:  $\delta$ 166.9, 168.8, 171.7, 172.3 and 173.8.

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#### Example 12

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HOOC-CH2-(ROIS) CH (COOH) - (R) Cha-hze-Pab/b x 2 HCl

# (i) BnOoC-CH<sub>2</sub>-(RorS) CH(COOBn)-(R) Cha-Aze-Pab(Z)/b

(R,S)CH(COOBn)-(R)Cha-Aze-Pab(Z). This diastereomer came The title compound was obtained by using the same procedure as described in Example 11 above on BnOOC-CH $_2$ out after the first one from the column. Yield 0.537 g. 20

## (ii) HOOC-CH<sub>2</sub>-(RorS) CH(COOH) - (R) Cha-Aze-Pab/b x 2 HCl 25

Removal of the catalyst by filtration and evaporation of the solvent in vacuo gave a residue which was dissolved in 6 ml water and 1.0 ml 1M hydrochloric acid. Freeze  $\operatorname{Pab}(2)/b$  dissolved in 15 ml 95% ethanol and 3 ml water was hydrogenated in the presence of 5% Pd/C for 5 h. 530 mg (0.65 mmol) BnOOC-CH<sub>2</sub>-(Rors)CH(COOBn)-(R)Cha-Azedrying gave 290 mg (78%) of the product.

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<sup>1</sup>H-NWR (300 MHz, MeOD, mixture of two rotamers): 6 0.86-1.90 (m, 13H), 2.30-2.42 (m, 1H), 2.60-2.75 (m,1H), 2.75-

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2.85 (m, 1H), 2.95-3.05 (m, 1H),3.65-3.71 (m, 1H), 4.00-4.10 (m, 1H), 4.14-4.24 (m, 1H),4.36-4.62 (m, 3H), 4.78-4.86 (m, 1H partially obscured by the solvent signal), 7.57 (d, 2H), 7.75 (d, 2H).

Resolved signals arising from a minor rotamer appears at 6 0.78, 2.92, 3.82, 5.36 and 7.80

 $^{13}\mathrm{C\text{-}NMR}$  (75 MHz  $\mathrm{D}_2\mathrm{O}$ ): amidine and carbonyl carbons: & 166.8, 169.0, 172.0, 172.4 and 175.2.

#### Example 13

## HOOC-CH2-CH2-(R)Cha-Aze-Pab x 2 HC1

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(i) BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cha-Aze-Pab(Z)

A mixture of 182 mg (0.35 mmol) H-(R)Cha-Aze-Pab(Z) (See Example 8) and 62.5 mg (0.385 mmol) benzyl acrylate in 1.5 ml 95% ethanol was stirred at room temperature for 4 days. The solvent was removed in vacuo and the residue was subjected to flash chromatography using ethyl acetate/methanol 9:1 as eluent to give 200 mg (84%) of the title compound.

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## (ii) HOOC-CH2-CH2-(R)Cha-Aze-Pab x 2 HC1

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195 mg (0.29 mmol) BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cha-Aze-Pab(Z) dissolved in 10 ml 95% ethanol and 2 ml water was hydrogenated in the presence of 5% Pd/C for 4 h. Removal of the catalyst by filtration and evaporation of the solvent in vacuo gave a residue which was dissolved in 2 ml water and 0.4 ml 1M hydrochloric acid. Freeze drying gave 130 mg (86%) of the product.

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<sup>1</sup>H-NMR (500 MHz,  $CD_3OD$ ): 6 0.98-1.27 (m, 2H), 1.30-1.90 (m, 11H), 2.27-2.35 (m, 1H), 2.65-2.74 (m, 1H), 2.77 (t,

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2H), 3.32 (t, 2H), 4.10 (t, 1H), 4.17-4.25 (m, 1H), 4.40-4.49 (m, 1H), 4.55 (AB, 2H), 4.83-4.90 (m, 1H), 7.58 (d, 2H), 7.77 (d, 2H).

 $^{13}\text{C-NMR}$  (125 MHz  $D_2\text{O}$ ): amidine and carbonyl carbons:  $\delta$  167.0, 168.9, 172.4 and 174.6.

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#### Example 14

# 10 HOOC-CH2-NH-CO-CH2-(R)Cha-Aze-Pab x 2 HC1

(i) BnOOC-CH<sub>2</sub>-NH-CO-CH<sub>2</sub>-(R)Cha-Aze-Pab(Z)

A mixture of 0.212 g ( 0.408 mmole ) H-(R)Cha-Aze-Pab(Z)(See Example 8), 0.124 g ( 0.89 mmole ) K<sub>2</sub>CO<sub>3</sub> and 0.128 g ( 0.449 mmole ) BnOOC-CH<sub>2</sub>-NH-CO-CH<sub>2</sub>-Br (See preparation of starting materials) in 6 ml acetonitrile was stirred at 50°C for two hours. After evaporation of the solvent the residue was dissolved in water and ethyl acetate. The water layer was extracted twice with ethyl acetate and the combined organic layer was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and evaporated. The product was purified by flash chromatography using a stepwise gradient of ethyl acetate/tetrahydrofurane ( 85/15, 4/1, 7/3 ) to yield 0.190 g ( 64% ) of the title compound.

1H-NMR (300 MHz, CDCl<sub>3</sub>): & 0.75-2.1 (m, 13H), 2.43 (m, 1H), 2.56 (d, 1H), 2.79 (m, 1H), 3.0-3.15 (m, 2H; thereof 3.05 (d, 1H)), 3.89-4.18 (m, 5H), 4.8-4.98 (m, 2H), 5.15 (s, 2H), 5.18 (s, 2H), 7.2-7.47 (m, 12H), 7.72 (t, NH), 7.86 (d, 2H), 8.14 (bs, NH), 8.31 (dd, NH), 9.42 (bs, NH)

<sup>13</sup>C-MMR (75 MHz, CDCl<sub>3</sub>): amidine and carbonyl carbons: 6 164.5, 168.7, 169.22, 169.83, 171.7, 175.5

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(ii) HOOC-CH2-NH-CO-CH2-(R)Cha-Aze-Pab x 2 HCl

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0.19 g ( 0.26 mmole ) of Bnooc-CH<sub>2</sub>-NH-CO-CH<sub>2</sub>-(R)Cha-Aze-pab(2) was mixed with 0.075 g \$ \$ Pd/C, 1.5 ml 1N HCl-solution, 3 ml water and 17 ml ethanol and the mixture was hydrogenated at atmospheric pressure for one hour. Filtration of the catalyst, evaporation of the solvent followed by freeze drying from water gave 144 mg (97 %) of the title compound.

<sup>1</sup>H-NMR ( D<sub>2</sub>O, 300 MHz, two rotamers 4:1): 6 0.88-1.88 (m, 13H), 2.25-2.42 (m, 1H), 2.63-2.89 (m, 1H), 3.94 (s, 2H), 3.99 (apparent doublet, 2H), 4.16 (t, 1H), 4.28 (q, 1H), 4.41 (q, 1H), 4.56 (s, 2H), 4.98 (dd, 1H), 7.53 (d, 2H), 7.71 (d, 2H),

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Resolved signals from the minor rotamer appears at 6 0.50 (bg), 0.77 (bg), 5.21 (dd), 7.56 (d) and 7.81 (d).

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13C-NMR ( D<sub>2</sub>O, 75 MHz ): The carbonyls and amidinecarbon at 6 166.8, 166.9, 168.6, 172.3 and 173.4.

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Resolved signals from the minor rotamer appears at 6: 166.6, 169.6 and 172.0

#### Example 15

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H-(R) Cha-Pro-Pab x 2 HCl

(i) Boc-(R)Cha-Pro-Pab(Z)

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0.135 ml (1.1 mmol) pivaloyl chloride was added at room temperature to a stirred mixture of 0.155 ml (1.1 mmol) triethyl amine and 405 mg (1.1 mmol) Boc-(R)Cha-Pro-OH triethyl amine and 405 mg (1.1 mmol) Boc-(R)Cha-Pro-OH After 3 h 340 mg (1.1 mmol) H-Pab(Z) (See preparation of starting materials) in 5 ml DMF was added and stirring was continued over night. The reaction mixture was diluted with water and extracted with ethyl

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acetate/toluene 1:1. The organic phase was dried (MgSO<sub>4</sub>) and the solvent was removed in vacuo to give a residue which was subjected to flash chromatography using ethyl acetate as eluent. The yield was 309 mg (49%).

(11) H-(R)Cha-Pro-Pab(Z)

Hydrogen chloride was bubbled through a solution, until saturation, of 1.246 g (2 mmol) Boc-(R)Cha-Pro-Pab(Z) in 20 ml ethyl acetate at room temperature. After 30 minutes sodium carbonate solution (10%) was added and the organic phase which separated was dried (K<sub>2</sub>CO<sub>3</sub>). The drying agent was washed with methylene chloride and the solvent was evaporated from the combined organic phases to give 1.11 g (100%) of the title compound.

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(iii) H-(R)Cha-Pro-Pab x 2 HCl

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ethanol was hydrogenated in the presence of 38 mg 10% pd/C for 1.5 h. Dilution of the reaction mixture with distilled water and removal of the catalyst by filtration followed by removal of the ethanol in vacuo and freeze drying gave the title compound as a colorless powder. The peptide was finally converted to the dihydrochloride by dissolution in hydrochloric acid followed by freeze drying to give 90 mg (100%) of the title compound.

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<sup>1</sup>H-NMR (300 MHz, D20); 6 1.0-2.0 (m, 13H), 2.0-2.3 (m, 3H), 2.3-2.5 (m, 1H), 3.6-3.7 (m, 1H), 3.8-3.9 (m, 1H), 4.5-4.6 (m, 3H), 7.4-7.6 (m, 3H), 7.6-7.9 (m, 2H).

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13C-NMR (75 MHz, D20): amidine and carbonyl carbons: 8 167.2, 170.0, 174.9.

## HOOC-CH2-(R)Cha-Pro-Pab x 2 HC1

(i) BnOOC-CH<sub>2</sub>-(R)Cha-Pro-Pab(Z)

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acetate as eluent to give 194 mg (57%) of the title which was subjected to flash chromatography using ethyl and the solvent was removed in vacuo to give a residue at 40°C for 2.5 h. The mixture was filtered through hyflo 181 mg (1.3 mmol)  ${
m K_2CO_3}$  in 2 ml acetonitrile was sonicated A mixture of 268 mg (0.5 mmol)  $H-\langle P \rangle$  Cha-Pro-Pab(Z) (See Example 15), 90  $\mu$ l (0.55 mmol) benzyl bromoacetate and

HOOC-CH2-(R)Cha-Pro-Pab x 2 HCl

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dried to give 115 (68%) of the desired product. was added and the resulting solution was finally freeze by freeze drying gave a white residue. Hydrochloric acid diluted with water and the catalyst was removed by mg 10% Pd on charcoal for 3 h. The reaction mixture was filtration. Evaporation of the ethanol in vacuo followed in 10 ml ethanol was hydrogenated in the presence of 77194 mg (0.28 mmol) Bn00C-CH<sub>2</sub>-(R)Cha-Pro-Pab(Z) dissolved

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3.8 (m, 1H), 3.8-4.0 (m, 3H), 4.4-4.7 (m, 4H), 7.5-7.7 1.5-2.0 (m, 8H), 2.0-2.3 (m, 3H), 2.3-2.5 (m, 1H), 3.6-(d, 2H), 7.7-7.9 (d, 2H). <sup>1</sup>H-NMR (300 MHz, D<sub>2</sub>0); & 1.0-1.2 (ш, 2H), 1.2-1.5 (ш, 3H),

167.1, 168.2, 169.3, 174.6.  $^{13}\mathrm{C\text{-}NMR}$  (75 MHz,  $\mathrm{D_20}$ ): amidine and carbonyl carbons: 6

### Example 17

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ROOC-CH2-(Me)(R)Cha-Pro-Pab

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## (i) Boc-(Me)(R)Cha-Pro-Pab(Z)

was stirred at room temperature for 2 days. to 8-9 with N-methylmorpholine, whereafter the solution DMF, and the pH of the resulting solution was adjusted Pab(2) (See preparation of starting materials) in 3 ml of DMF was added a solution of 0.562 g (1.85 mmol) of H-Pro-OSu (See preparation of starting materials) in 3 ml To a solution of 0.8 g (1.67 mmol) of Boc-(Me)(R)Cha-

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compound as a yellowish white powder. Evaporation of the solvent gave 0.65 g (60%) of the title  ${\tt NaHCO_3}$  solution, water and brine, and dried  $({\tt Na_2SO_4})$ . organic solution was washed with 1M KHSO $_{
m 4}$  solution, 10 ${
m \$}$ mixture was extracted with 3x25 ml of ethyl acetate. The The solution was poured onto water, and the resulting

## (ii) Me-(R)Cha-Pro-Pab(Z)

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25 20 white fluffy powder. and evaporated to give 0.4 g (79%) of the compound as a 3x25 ml ethyl acetate. The extract was washed with brine residue was dissolved in a  $ext{Na}_2 ext{CO}_3$  solution, extracted with The resulting solution was evaporated to dryness, and the and the solution was stored in refrigerator overnight. Pab(Z) in 50 ml of EtoH was saturated with HCl at 0°C, A solution of 0.60 g (0.92 mmol) of Boc-(Me)(R)Cha-Pro-

30 4.61 (d, 1H), 5.2 (s, 2H), 7.2-7.45 (m, 7H), 7.72 (t, 3.28 (dd, 1H), 3.41 (q, 1H), 3.62 (m, 1H), 4.42 (m, 2H), <sup>1</sup>H-NMR (CDCl<sub>3</sub>, 300 MHz): δ 0.8-1.0 (m, 2H), 1.1-1.4 (m, 5H), 1.4-1.55 (m, 1H), 1.6-1.9 (m, 10H), 1.9-2.05 (m, 1H), 7.79 (d, 2H). 2.05-2.2 (m, 2H), 2,19 (s,3H), 2.4-2.5 (m, 1H),

# (iii) Bn00C-CH2-(Me)(R)Cha-Pro-Pab(Z)

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A mixture of 0.40 g (0.73 mmol) of Me-(R)Cha-Pro-Pab( $\mathbb{Z}$ ),

evaporated. The crude product (0.69 g) was subjected to flash chromatography (CH2Cl2/MeOH 10/1) yielding 0.39 g temperature overnight. The resulting mixture Was evaporated, ethyl acetate was added, and the mixture was washed with water and brine, dried (Na2SO4), and (mortared) in 15 ml of CH<sub>3</sub>CN was stirred at room 0.17 g Bnooc-CH<sub>2</sub>Br and 0.20 g (2 equiv.) of  $K_2^{\text{CO}_3}$ (77%) of a light yellow very viscous oil.

HOOC-CH2-(Me) (R) Cha-Pro-Pab 20

freeze dried to yield 0.25 g (95%) of the compund as a (Me) (R) Cha-Pro-Pab(Z) in 30 ml of EtOH was added 0.1 g of Pd/C (10%), and the substance was hydrogenated at atmospheric pressure. The solution was filtered and evaporated, whereafter the remaining syrupy material was To a solution of 0.39 g (0.56 mmol) of Bnooc-CH2white crystalline powder.

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3.75 (m, 2H), 3.95-4.1 (m, 2H), 4.35-4.5 (m, 3H), 7.55 1H), 2.15-2.3 (m, 1H), 2.57 (s, 3H), 3.32 (d, 1H), 3.55-<sup>1</sup>H-NMR (300 MHz, CD<sub>3</sub>OD): 6 0.85-1.1 (m, 2H), 1.1-1.4 (m, 6H), 1.5-1.85 (m, 9H), 1.9-2.05 (m, 3H), 2.05-2.15 (m, (d, 2H), 7.72 (d, 2H).

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 $^{13}\mathrm{C-NMR}$  (75 MHz,  $^{\mathrm{CD}_3\mathrm{OD}}$ ): amidine and carbonyl carbons: \$ 168.4, 171.5, 174.7, 175.1.

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Example 18

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HOOC-CH2-CH2-(R) Cha-Pro-Pab x 2 HCl

(i) Bnooc-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cha-Pro-Pab(Z)

Example 15) and 66 mg (0.4 mmol) benzyl acrylate in 1.5 A mixture of 149 mg (0.28 mmol) H-(R)Cha-Pro-Pab(Z) (See ml ethanol was kept at room temperature for 36 h. The 35

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solvent was removed in vacuo and the residue was subjected to flash chromatography using ethyl acetate as eluent to give 124 mg (64%) of the desired product.

(ii) HOOC-CH2-CH2-(R)Cha-Pro-Pab x 2 HCl ß

residue was dissolved in hydrochloric acid and the dissolved in 10 ml ethanol was hydrogenated for 1 h in by filtration and the solvent was removed in vacuo. The resulting solution was freeze dried to give 87 mg (79%) the presence of 55 mg 10% Pd/C. The catalyst was removed (0.18 mmol) BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cha-Pro-Pab(Z) of the title compound. 124 mg

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3.3-3.4 (m, 1H), 3.5-3.7 (m, 1H), 3.7-3.9 (m, 1H), 4.3-3H), 2.2-2.4 (m, 1H), 2.7-2.8 (t, 2H), 3.2-3.3 (m, 1H), H-NWR (300 MHz, D20): 6 1.0-2.0 (m, 13H), 2.0-2.2 (m, 4.6 (m, 4H), 7.4-7.6 (m, 2H), 7.7.6-7.8 (m, 2H). 12

 $^{13}\text{C-NMR}$  (75 MHz,  $^{12}\text{O}$ ): amidine and carbonyl carbons:  $^{6}$ 167.0, 168.3 and 174.6 (Two carbons are overlapping). 20

Example 19

HOOC-CH2-CH2-(Me) (R) Cha-Pro-Pab x 2 HC1 25 (i) BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(Me)(R)Cha-Pro-Pab(Z)

amount of 16.2 mg (0.1 mmol) of benzyl acrylate was added and the stirring continued for 24 h. The solvent was evaporated and the residue was subjected to flash chromatography ( $\mathrm{CH_2Cl_2/MeOH(NH_3-saturated)}$ , 95/5) to give 97.3 mg (0.6 mmol) of benzyl acrylate and the reaction was stirred at room temperature. After 72 h an additional Pab(Z) (See Example 17) in 5 ml of EtOH (99%) was added To a solution of 274 mg (0.5 mmol) of Me-(R)Cha-Pro-198 mg (56%) of the title compound. 35 30

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13C-NMR (125 MHz, CDCl<sub>3</sub>): amidine and carbonyl carbons: 6 164.7, 167.9, 171.7, 172.3 and 172.6.

# (ii) $HOOC-CH_2-CH_2-(Me)(R)Cha-Pro-Pab x 2 HC1$

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To a solution of 198 mg (0.27 mmol) BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(Me)(R)Cha-Pro-Pab(Z) in 10 ml EtOH and 1 ml 1M HCl was added 60 mg of 5 % Pd/C (containg 50 % H<sub>2</sub>O by weight) and the mixture was hydrogenated at athmospheric pressure for 4 h. The catalyst was filtered off and the solvent was evaporated. The remaining oil was dissolved in water and freeze dried to give the title compound in a quantitative yield.

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1H-NMR (500 MHz, D<sub>2</sub>O): \$ 1.08-1.2 (m, 2H), 1.2-1.42 (m, 4H), 1.68-1.91 (m, 5H), 1.93-2.08 (m, 2H), 2.09-2.26 (m, 3H), 2.49 (m, 1H), 2.95 (m, 2H), 3.03 (s, 3H), 3.60 (apparent bs, 2H), 3.82 (m, 1H), 3.98 (m, 1H), 4.53 (m, 1H), 4.61 (bs, 2H), 4.64 (m, 1H), 7.63 (d, 2H), 7.97 (d, 2H).

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 $^{13}\text{C-NMR}$  (75 MHz,  $D_2\text{O}$ ): amidine and carbonyl carbons:  $\delta$  167.2, 167.8 and 174.5. Two peaks are probably overlapping.

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#### Example 20

# HOOC-CH2-(Rors)CH(COOH)-(R)Cha-Pro-Pab/a x 2 HCl

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(i) Bn00C-CH<sub>2</sub>-(R,S)CH(COOBn)-(R)Cha-Pro-Pab(Z)

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A mixture of 0.50 g (0.94 mmol) of H-(R)Cha-Pro-Pab(Z) (See Example 15) and 0.28 g (0.94 mmol) of dibenzyl maleate in 20 ml of EtOH was kept at room temperature for 5 days. Evaporation of the solvent followed by flash chromatography using CH<sub>2</sub>Cl<sub>2</sub>/MeOH as eluent gave 0.15 g (19

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\$) of the diastereomeric mixture.

1H NMR (500 MHz, CDCl<sub>3</sub>) & 0.7-2.1 (m, 17 H), 2.3-2.4 (m,
1 H), 2.5-2.8 (m, 2 H), 3.2-3.7 (m, 4 H), 4.46 (d, 1 H),
4.65 (bd, 1 H), 4.81 (d, 1 H), 4.9-5.1 (m, 3 H), 5.20 (s,
2 H), 7.1- 7.4 (m, 15 H), 7.4-7.5 (m, 2 H), 7.6-7.8 (m,
3 H).

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# (ii) HOOC-CH<sub>2</sub>-(Rors)CH(COOH)-(R)Cha-Pro-Pab/a x 2 HCl

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A mixture of 0.15 g (0.18 mmol) of Bnooc-CH<sub>2</sub>-(R,S)CH(COOBn)-(R)Cha-Pro-Pab(Z) was dissolved in 5 ml of ethanol and was hydrogenated over 5% Pd/C at atmospheric pressure for 1 h. to give HOOC-CH<sub>2</sub>-(R,S)CH(COOH)-(R)Cha-Pro-Pab.The two diastereomers were separated by RPJC using (CH<sub>3</sub>CN/0.1 M NH<sub>4</sub>OAC 15/85) as eluent followed by freeze drying from HCl. This diastereomer eluted first from the column. Yield 19 mg (18 %).

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1H-NMR (500 MHz, D<sub>2</sub>O, mixture of two rotamers) major rotamer: 6 1.0-2.0 (m, 15H), 2.15 (m, 2H), 2.44 (m, 1H), 3.00 (bd, 1H), 3.05 (bd, 1 H), 3.69 (m, 1H), 3.84 (m, 1H), 3.97 (bs, 1H), 4.5-4.7 (m, 3H), 7.62 (d, 2H), 7.87 (d, 2H).

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13C-NMR (75 MHz, D<sub>2</sub>0): amidine and carbonyl carbons: & 167.2, 168.3, 173.8, 174.6 and 178.2.

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#### Example 21

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HOOC-CH2-(Rors) CH (COOH)-(R) Cha-Pro-Pab/b x 2 HC1

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Constitution of ي

(R,S)CH(COOH)-(R)Cha-Pro-Pab. This diastereomer came out The title compound was obtained by using the same after the first one from the column. Yield 19 mg (18 %). procedure as descibed in Example 20 on HOOC-CH $_2$ -  $^{1} ext{H-NMR}$  (500 MHz,  $ext{D}_{2} ext{O}$ , mixture of two rotamers) major 1H), 4.03 (bs, 1H), 4.5-4.7 (m, 3H), 7.58 (d, 2H), 7.84 rotamer: 6 1.0-2.0 (m, 14H), 2.15-2.25 (m, 3H), 2.44 (m, 1H), 3.11 (bd, 1H), 3.19 (bd, 1H), 3.71 (m, 1H), 3.92 (m, (d, 2H). Resolved signals arising from the minor rotamer appears at: 6 7.66 (d) and 7.91 (d).

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 $^{13}\mathrm{C-NMR}$  (75 MHz,  $^{13}\mathrm{D}_2\mathrm{O}$ ): amidine and carbonyl carbons:  $^{\delta}$ 167.3, 168.5 and 174.7. Two carbons are probably overlapping. 12

#### Example 22

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HOOC-CH2-NH-CO-CH2-(R)Cha-Pro-Pab x 2 HCl

(i) BnOOC-CH<sub>2</sub>-NH-CO-CH<sub>2</sub>-(R)Cha-Pro-Pab(Z)

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water layer was extracted one more time with ethyl filtered and evaporated to yield 0.350 g of an oil. The crude product was purified by flash chromatography using a stepwise gradient of  $\mathrm{CH_2Cl_2/MeOH~97/3},~95/5,~92.5/7.5$  to was stirred at 50°C for 2 h 30 minutes, the solvent was evaporated and the residue was partitioned between water and ethyl acetate. The layers were separated and the  $_{\mathrm{mmole}})$  BnOOC-CH $_{2}$ -NH-CO-CH $_{2}$ -Br (See preparation of starting materials) was mixed in 6 ml acetonitrile. The mixture acetate. The combined organic layer was dried (  ${
m Na}_2{
m SO}_4$ ), Example 15), 0.140 g (1.01 mmole)  $K_2\mathsf{CO}_3$  and 0.145 g (0.506 0.246 g (0.460 mmole) of H-(R)Cha-Pro-Pab(Z) yield 0.227 g (67%) of the title compound.

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13C-NMR (75 MHz, CDCl3): 6 25.0, 26.0, 26.2, 26.4, 26.7, 32.4, 34.2, 34.4, 40.8, 40.9, 42.9, 46.7, 50.5, 58.4, 60.2, 67.0, 67.2, 127.5, 127.8, 128.2, 128.3, 128.4, 128.5, 128.6, 128.6, 134.1, 135.2, 137.0, 142.6, 164.7, 168.9, 169.3, 170.4, 172.2, 175.0

(11) HOOC-CH2-NH-CO-CH2-(R)Cha-Pro-Pab x 2 HCl

pressure for one and a half hour. Filtration of the catalyst through hyflo and freeze drying with  $lml\ lN$ hydrochloric acid gave 0.058 g (82%) of the desired acetic acid. The mixture was hydrogenated at athmospheric was mixed with 30 mg 5 % Pd/C and dissolved in 10 ml 0.089 g (0.12 mmole) BnOOC-CH<sub>2</sub>-NH-CO-CH<sub>2</sub>-(R) Cha-Pro-Pab(Z) 12 10

1H), 3.55-3.7 (m, 1H), 3.7-4.1 (m, 5H),4.42 (t, 1H),  $^{1}\mathrm{H-NMR}$  (300 MHz, D<sub>2</sub>0): 6 0.9-2.2 (m, 16H), 2.25-2.47 (m, 4.48-4.6 (m, 3H), 7.51 (d, 2H), 7.77 (d, 2H)

20

 $^{12}\mathrm{C-NMR}$  (75 MHz, D<sub>2</sub>0): amidine and carbonyl carbons:  $^{6}$ 166.8, 167.1, 168.2, 173.6 and 174.6

22

Example 23

Etcoc-CH2-CH2-CH2-(R) Cha-Pro-Pab x HOAc

(i) EtOOC-CH=CH-CH<sub>2</sub>-(R)Cha-Pro-Pab(2) 30

BrCH2CH=CHCOOEt (108 mg, 0.56 mmol) in CH3CN (10 ml) at residue was dissolved in EtOAc (5 ml)/ $H_2$ O (2 ml). The mmol) was treated with  $\rm K_2CO_3$  (141 mg, 1.02 mmol) and H-(R)Cha-Pro-Pab(2) (See Example 15) (275 mg, 0.51 20°C for 20 h. The solvent was evaporated and the organic layer was separated, dried (Na<sub>2</sub>SO<sub>4</sub>), and

concentrated yielding 397 mg of an oil which was purified by flash chromatography using EtoAc/Heptane, 1/4 as eluent to give 252 mg (77%) of the title compound.

1H-NMR (500 MHz, CDCl<sub>3</sub>): & 0.8-1.05 (m, 2H), 1.1-1.45 (m, 3H), 1.3 (t, 3H), 1.5-1.9 (m, 8H), 1.95-2.05 (m, 1H), 2.1-2.15 (m, 1H), 2.45-2.55 (m, 1H), 3.0 and 3.15 (two d, 2H), 3.35-3.45 (m, 2H), 3.55-3.65 (m, 1H), 4.15 (q, 2H), 4.3 (d, 1H), 4.6-4.7 (m, 2H), 5.2 (s,2H), 5.85 (d, 1H), 6.75 (dt, 1H), 5.3-5.4 (m, 4H), 7.45 (d, 2H), 7.85 (d, 2H),

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13C-NMR (75.0 MHz, CDCl<sub>3</sub>): amidine and carbonyl carbons: 8 165.7, 171.2 and 175.7 (two peaks are probably overlapping).

(ii) EtOOC-CH2-CH2-CH2-(R)Cha-Pro-Pab x HOAc

20

EtOOCCH=CHCH<sub>2</sub>-(R)Cha-Pro-Pab(Z) (250 mg, 0.38 mmol) was disolved in ethanol and hydrogenated in the presence of 5 % Pd/C during approximately 2 h. Removal of the catalyst by filtration and evaporation of the solvent in vacuo gave after purification by RPLC using (CH<sub>3</sub>CN/0.1 M NH<sub>4</sub>OAc) as eluent 70 mg (36%) of the desired product.

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<sup>1</sup>H NMR (500 MHz, CD<sub>3</sub>OD):  $\delta$  0.9-1.05 (m, 2H), 1.15-1.55 (m, 5H), 1.25 (t, 3H), 1.6-1.85 (m, 7H), 1.95-2.6 (m, 8H), 3.55-3.65 (m, 2H), 3.8 (m, 1H), 4.1 (q, 2H), 4.45 (m and d, 2H), 4.55 (d, 1H), 7.55 and 7.75 (two d, 4H).

<sup>13</sup>C-NMR (75.0 MHz, CD<sub>3</sub>OD): amidine and carbonyl carbons: 6 168.3, 173.2, 174.6 and 174.9.

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Example 24

Ph(4-COOH)-802-(R)Cha-Pro-Pab x HC1

5 (i) Ph(4-COOH)-SO<sub>2</sub>-(R)Cha-Pro-Pab(Z)

15 10 chloride/methanol 3:1 as eluents gave 82 mg (39%) of the acetate/methanol of the residue by flash chromatography using ethyl product. and after 24 hours it was washed with water and dried The mixture was slowly allowed to reach room temperature  $(\mathrm{Na_2SO_4})$  . Removal of the solvent in vacuo and purification (0.58 mmol) triethyl amine in 4 ml methylene chloride. mmol)  $H^-(R)$  Cha-Pro-Pab(Z) (See Example 15) and 59 mg at ice bath temperature to a solution of 156 mg (0.29 64 mg (0.32 mmol) 4-chlorosulfonyl-benzoic acid was added 9:1 followed methylene

(ii) Ph(4-COOH)-SO2-(R)Cha-Pro-Pab x HC1

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80 mg (0.11 mmol) Ph(4-COOH)-SO<sub>2</sub>-(R)Cha-Pro-Pab(Z) was hydrogenated over 5 % Pd/C in EtOH. The catalyst was filtered off, the solvent evaporated and the crude product was purified by RPIC using (CH<sub>2</sub>CN/0.1 M NH<sub>4</sub>OAc 1/4) as eluent and finally converted to the hydrochloride salt by freeze drying from HCl which gave 21 mg (29%) of the product.

35  $^{13}\text{C-NMR}$  (75 MHz, CD<sub>3</sub>OD): amidine and carbonyl carbons:  $\ell$  168.4, 173.4, 173.9 and 174.2

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MS m/z 584 (M+ +1)

Example 25

H-(R)Cha-Pic-Pab x 2 HCl

(i) Boc-(R)Cha-Pic-Pab(Z)

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graphy on silica gel using ethyl acetate/toluene 2:1 as starting materials), 9.07 g (74.2 mmol) DMAP and 5.26 g (18.6 mmol) H-Pab(Z) (See preparation of starting naterials) in 200 ml DMF. The temperature was allowed to rise to 20°C over night. The solvent was removed in vacuo and toluene and water was added. The organic phase was washed with water, 1M KHSO4, 10% Na2CO3 and brine. Drying  $(\mathrm{MgSO}_4)$  and evaporation of the solvent in vacuo gave 13.63 3.57 g (18.6 mmol) EDC was added at -15°C to a mixture of 7.11 g (18.6 mmol) Boc-(R)Cha-Pic-OH (See preparation of g of a residue which was subjected to flash chromatoeluent to give 9.5 g (79%) of the title compound.

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25H), 2.3-2.5 (m, 1H), 2.9-3.1 (m, 1H), 3.8 (d, 1H), 4.3 (dd, 1H), 4.4-4.6 (m, 2H), 5.1 (s, 2H), 5.1-5.3 (m, 2H), 7.2-7.3 (m, 5H), 7.35 (d, 2H), 7.4-7.5 (m, 1H), 7.75 (d, <sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>): 6 0.7-1.0 (m, 2H), 1.0-2.2 (m,

25

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13C-NMR (75 MHz, CDCl3): amidine and carbonyl carbons: 6 156.8, 164.6, 168.2, 170.0 and 173.4.

(11) H-(R)Cha-Pic-Pab(Z)

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Aydrogen chloride was bubbled through a solution of 9.5 g (14.7 mmol) Boc-(R)Cha-Pic-Pab(Z) in 100 ml ethyl acetate at room temperature until saturation. After 10 minutes  $\mathrm{Na_2CO_3}$  solution (10%) was added and the organic phase which separated was dried  $(K_2\text{CO}_3)$  and the solvent

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was removed in vacuo to give the title compound in quantitative yield.

H-NMR (500 MHz, CD<sub>3</sub>OD): 6 0.85-1.05 (m, 2H), 1.15-1.90 (m, 16H), 2.25-2.35 (m, 1H), 3.20-3.30 (m, 1H), 3.80-3.90 (d, 1H), 3.90-4.0 (m, 1H), 4.4-4.5 (two d, 2H), 4.7 (br s, 5H) 5.15 (s, 2H), 5.20 (m, 1H), 7.25-7.45 (m, 7H), 7.85 (d, 2H).

(iii) H-(R)Cha-Pic-Pab x 2 HCl ព

mixture of 5 ml ethanol and 0.45 ml 1M hydrochloric acid was hydrogenated in the presence of 33 mg 10% Pd/C for 1.5 h. Removal of the catalyst by filtration and evaporation of the solvent in vacuo gave a residue which The purified peptide was finally converted to the dihydrochloride salt by dissolution in hydrochloric acid followed by freeze drying. The yield was 17 mg (35%) of was subjected to RPLC using 0.1 M  ${
m NH_4OAc/CH_3CN}$  as eluent. 55 mg (0.1 mmol) H-(R)Cha-Pic-Pab(Z) dissolved in the title compound

12

H-NMR (300 MHz,  $\mathrm{D_2O}$ , 2 rotamers, 3:1 mixture): 6 1.0-2.0 (m, 18H), 2.33 (d, 1H), 3.4-3.5 (m, 1H), 3.8-3.9 (m, 1H), 4.4-4.8 (m, 3H), 5.15.5.25 (m, 1H), 7.5-7.7 (m, 2H), 7.8-8.0 (m, 2H).

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Resolved signals from the minor rotamer appears at 0.5-0.7 (m) and 3.0-3.1 (m) 13C-NMR (75 MHz, D20): amidine and carbonyl carbons: 167.3, 171.6 and 173.6.

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Resolved signals for the minor rotamer appears at 6 170.6 and 172.4. 35

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Example 26

HOOC-CH2-(R)Cha-Pic-Pab x 2 HC1

(i) BnOOC-CH2-(R)Cha-Pic-Pab(Z)

mg (77%) of the desired product. residue was subjected to flash chromatography to give 720 at 40°C for 40 minutes. The solvent was removed and the 558 mg (4 mmol)  $\mathrm{K}_2\mathrm{CO}_3$  in 4 ml acetonitrile was sonicated Example 25), 230 ml (1.45 mmol) benzyl bromoacetate and A mixture of 742 mg (1.35 mmol) H-(R)Cha-Pic-Pab(Z) (See

15 1H), 4.35 (dd, 1H), 4.55 (dd, 1H), 4.80 (two d, 2H), 5.2 3.25 (d, 1H), 3.45 (d, 1H), 3.55-3.65 (m, 1H), 3.7 (m, 16H); 2.1-2.4 (br s, 1 or 2H), 2.4 (d, 1H), 3.0 (m, 1H), <sup>1</sup>H-NMR (500 MHz, CDCl<sub>3</sub>); & 0.8-1.0 (m, 2H), 1.1-1.9 (m, 2H), 5.3 (m, 1H), 7.2-7.4 (m, 12H), 7.8 (d, 2H).

20 164.5, 167.9, 170.5, 173.4 and 175.5.  $^{13}\mathrm{C\text{-}NMR}$  (125 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $\delta$ 

(ii) HOOC-CH<sub>2</sub>-(R)Cha-Pic-Pab x 2 HCl

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freeze dried to give 281 mg (79%) of the title compound. Hydrochloric acid was added and the solution was finally a residue which was dissolved in distilled water. filtration and evaporation of the solvent in vacuo gave mg 10% Pd/C for 4 h. Removal of the catalyst by 509 mg (0.73 mmol) BnOOC-CH $_2$ -(R)Cha-Pic-Pab(Z) dissolved in 25 ml ethanol was hydrogenated in the presence of 259

rotamer: & 1.0-2.0 (m, 18H), 2.25-2.40 (m, 1H), 3.4-3.5 (m, 1H), 7.55-7.75 (m, 2H), 7.8-8.0 (m, 2H). (m, 1H), 3.8-3.95 (m, 3H), 4.55-4.65 (two d, 2H), 5.15  $^{1}\mathrm{H^{-}NMR}$  (500 MHz, D<sub>2</sub>O, mixture of rotamers 4:1): major

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Resolved signal for the minor rotamer appears at & 166.9, 167.3, 169.9, 170.3 and 173.5.  $^{13}\mathrm{C-NMR}$  (125 MHz, D $_2$ O): amidine and carbonyl carbons:  $\delta$ 

169.2 and 172.0

Example 27

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HOOC-CH2-(Rors)CH(COOH)-(R)Cha-Pic-Pab/a x 2 HCl

(i) BnOOC-CH<sub>2</sub>-(R,S)CH(COOBn)-(R)Cha-Pic-Pab(Z)

10

methanol/methylene chloride as eluent to give 275 mg solvent was removed in vacuo and the residue was ml ethanol was kept at room temperature for 1 week. The (30%) the diastereomeric mixture. subjected Example 25) and 332 mg (1.1 mmol) dibenzyl maleate in 1 A mixture of 592 mg (1.1 mmol) H-(R)Cha-Pic-Pab(Z) (See ţ flash chromatography

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(ii) HOOC-CH<sub>2</sub>-(RorS)CH(COOH)-(R)Cha-Pic-Pab/a x 2 HCl

25 5 This diastereomer eluted first from the column. Yield 9  $\mathrm{NH_{4}OAc}$  1/4 ) as eluent followed by freeze drying from HCl. diastereomers were separated by RPLC using  $(CH_3CN/0.1\ M_3CN/0.1\ M_3CN/0.$ 166 mg of HOOC-CH2-(R,S)CH(COOH)-(R)Cha-Pic-Pab. The two vacuo. Addition of water followed by freeze drying gave hours in the presence of 75 mg 10% Pd/C. The mixture was dissolved in 20 ml 95% ethanol was hydrogenated for 18 filtered through hyflo and the solvent was removed in  $BnOOC-CH_2-(R,S)CH(COOBn)-(R)Cha-Pic-Pab(Z)$ 

30

18H), 2.25-2.4 (m, 1H), 3.0-3.2 (m, 2H), 3.4 (t, 1H), 3.8 (d, 2H), 7.9 (d, 2H). (d, 1H), 4.05 (t, 1H), 4.5-4.7 (m, 3H), 5.2 (s, 1H), 7.55  $^{
m 1}$ H-NMR (300 MHz, D $_{
m 2}$ O, mixture of rotamers): & 1.0-2.0 (m,

<u>ა</u>

LC1

Resolved signals from the minor rotamer appears at 4.0(t) and 7.7(d).

#### Example 28

# $\mathtt{HOOC-CH_2-(ROIS)CH(COOH)-(R)Cha-Pic-Pab/b \ x \ 2 \ HCl}$

The title compound was obtained by using the same procedure as described in Example 27 on HOOC-CH<sub>2</sub>-(R,S)CH(COOH)-(R)Cha-Pic-Pab. This diastereomer came out after the first one from the column.

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1H-NMR (500 MHz, D20, mixture of rotamers; 6 1.0-2.0 (m, 18H), 2.25-2.4 (m, 1H), 3.0-3.2 (m, 2H), 3.5 (t, 1H), 3.85 (d, 1H), 4.15 (s, 1H), 4.5-4.7 (m, 3H), 5.15 (s, 1H), 7.55 (d, 2H), 7.8 (d, 2H).

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Resolved signals from the minor rotamer appear at  $\delta$  4.35(s), 7.65(d) and 7.9(d).

#### Example 29

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 $HOOC-CH_2-CH_2-(R)$  Cha-Pic-Pab x 2 HCl

25 (i) BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cha-Pic-Pab(Z)

A mixture of 851 mg (1.55 mmol) H-(R)Cha-Pic-Pab(2) (See Example 25) and 269 mg (1.71 mmol) benzyl acrylate in 5 ml ethanol was kept at room temperature for 40 h. The solvent was removed in vacuo and the residue was subjected to flash chromatography using methylene chloride/methanol as eluent to give 812 mg (74%) of the product.

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1H-NMR (500 MHz, CDCl<sub>3</sub>): 6 0.8-1.0 (m, 2H), 1.1-1.9 (m, 16H), 2.3-2.5 (m, 3H), 2.6-2.8 (m, 2H), 3.0 (m, 1H), 3.5 (m, 1H), 3.5-3.7 (m, 1H), 4.3 (dd, 1H), 4.6 (dd, 1H),

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4.95-5.05 (two d, 2H), 5.2 (s, 2H), 5.3 (m, 1H), 6.5-6.9 (br s, 1H), 7.0-7.1 (m, 1H), 7.2-7.5 (m, 12H), 7.75-7.85 (d, 2H), 9.3-9.7 (br s, 1H).

# (ii) $HOOC-CH_2-CH_2-(R)Cha-Pic-Pab \times 2 HCI$

dissolved in 25 ml ethanol was hydrogenated for 4 h in dissolved in 25 ml ethanol was hydrogenated for 4 h in the presence of 306 mg 15% Pd/C. The catalyst was removed by filtration and the solvent was removed in vacuo. The residue was dissolved in hydrochloric acid and the resulting solution was freeze dried to give 481 mg (78%) of the title copound.

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15 H-NMR (500 MHz, D<sub>2</sub>0): 6 0.95-1.1 (m, 2H), 1.15-1.9 (m, 16H), 2.2-2.3 (m, 1H), 2.7-2.8 (t, 2H), 3.2-3.3 (m, 3H), 3.4-3.5 (m, 1H), 3.75-3.85 (m, 1H), 4.4-4.6 (m, 3H), 5.15 (m, 1H), 7.5-7.6 (m, 2H), 7.8-7.9 (m, 2H), 8.6-8.7 (m, 1H)

13C-NWR (125 MHz, CD<sub>3</sub>OD): amidine and carbonyl carbons: *§* 170.6, 175.9, 179.5 and 183.5.

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#### Example 30

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HOOC-CO-(R) Cha-Pic-Pab x HOAG

## (i) EtOOC-CO-(R)Cha-Pic-Pab(Z)

0.12 g ethyloxalyl chloride was added to a mixture of 0.42 g (0.77 mmol) H-(R)Cha-Pic-Pab(2) (See Example 25) and 0.21 g (1.5 mmol) K<sub>2</sub>CO<sub>3</sub> in 10 ml CH<sub>3</sub>CN at room temperature. After 2 hours an additional amount of 0.07 g (0.5 mmol) ethyloxalyl chloride was added. The mixture was stirred at room temperature over night. The solvent was removed in vacuo. and the residue was dissolved in CH<sub>2</sub>Cl<sub>2</sub> and washed with water. Evaporation and flash

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## (11) HOOC-CO-(R)Cha-Pic-Pab(Z)

CH2Cl2: methanol) gave 0.21 g (42%) of the product.

evaporated to give 80 mg of the product. The aqueous phase was acidified with 0.5M HCl (pH 1) onto ethyl acetate/water. The phases were separated and and extracted with  $ext{CH}_2 ext{Cl}_2$ , dried over  $ext{Na}_2 ext{SO}_4$  and the organic phase was extracted with a KHCO3-solution. stirred at room temperature over night and then poured dissolved in 3 ml water was added. The mixture was dissolved in 3 ml THF and 0.17 g (4.2 mmol) LiOH 0.21 g (0.32 mmol) EtOOC-CO-(R)Cha-Pic-Pab(Z) was

## (iii) HOOC-CO-(R)Cha-Pic-Pab x HOAc

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RPLC to give the title compound. evaporated. The residue was subjected to purification by in EtOH. The catalyst was filtered off and the solvent HOOC-CO-(R)Cha-Pic-Pab(Z) was hydrogenated over 5 % Pd/C

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by the solvent signal. 8.22-8.27 (m, 1H), 9.32 (broad s), 9.90 (broad s). The signal of one of the protons (3.25) is partially obscured (m, 1H), 7.41 (d, 2H), 7.75 (d, 2H), 8.1-8.15 (m, 1H), 1H), 4.32, 4.44 (AB, 2H), 4.71-4.77 (m, 1H), 4.98-5.02 15H), 1.86-1.94 (m, iH), 2.13-2.2 (m, 1H), 3.75-3.81 (m, <sup>1</sup>H NMR (500 MHz, DMSO-d<sub>6</sub>); & 0.8-1.0 (m, 2H), 1.1-1.75 (m,

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MS m/z 486 (M+ + 1)

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#### Example 31

## HOOC-CH2-CO-(R) Cha-Pic-Pab

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## (1) $Meooc-CH_2-CO-(R)Cha-Pic-Pab(Z)$

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eluent gave 0.27 g (58%) of the desired product. dried (NaSO $_4$ ). Evaporation of the solvent followed by flash chromatography using toluen/ethyl acetate (1/3) as precipitated DCU was removed by filtration and the filtrate was washed with 0.3M KHSO $_{
m 4}$  and KHCO $_{
m 3}$ -solution and solution was stirred in room temperature over night. The 40 ml  $\mathrm{CH_2Cl_2}$  and 0.16 g (0.8 mmol) DCC was added. The and 0.9 g (0.8 mmol) monomethylmalonate was dissolved in 0.39 g (0.72 mmol) H-(R)Cha-Pic-Pab(Z) (See Example 25)

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## (ii) MeOOC-CH2-CO-(R)Cha-Pic-Pab

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15 by filtration and evaporation of the solvent gave 50  ${ t mg}$ presence of 5% Pd/C for 5 hours. Removal of the catalyst dissolved in 10 ml ethanol and was hydrogenated in 90 mg (0.14 mmol) MeOOC-CH2-CO-(R)Cha-Pic-Pab(Z) was (70%) of the title product.

- 20 3.95-4.05 (m, 1H), 4.4-4.55 (m, 3H), 5.15-5.25 (m, 1H), 7.4-7.55 (m, 2H), 7.7-7.85 (m, 2H). 16H), 2.35-2.45 (m, 1H), 3.2-3.4 (m, 3H), 3.7 (s, 3H), <sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD): 6 0.85-1.1 (m, 2H), 1.1-1.9 (m,
- 25 δ 168.2, 168.7, 170.0, 172.4 and 174.6. <sup>13</sup>C NMR (75 MHz, CD<sub>3</sub>OD): amidine and carbonyl carbons:

MS m/z 514 (M+ + 1)

### 30 (iii) HOOC-CH2-CO-(R)Cha-Pic-Pab

was extracted out from the insoluble inorganic salts with The aqueous phase was freeze dried. The soluble material water was added and the methanol was removed in vacuo. NaOH at room temperature. After stirring for 5 hours (R)Cha-Pic-Pab in 5 ml methanol was added 2 ml of 0.5 M To a solution of 0.14 g (0.27 mmol) of MeOOC-CH $_2$ -CO-

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absolute ethanol. The remaining solid after evaporation of the ethanol was suspended in water and 70 mg (52%) of the title compound was isolated by filtration.

16H), 2.15-2.30 (m, 1H), 2.58, 2.86 (AB, 2H), 3.8-3.95 (m, 1H), 9.90 (broad s, 3H). The signal of one of the 1H), 7.40 (d, 2H), 7.77 (d, 2H), 8.2-8.3 (m, 1H), 9.3-9.4 (m, 1H), 4.2-4.5 (m, 2H), 4.7-4.85 (m, 1H), 4.95-5.05 (m, protons (3.21) is partially obscured by the solvent-<sup>1</sup>H NMR (300 MHz, DMSO-d<sub>6</sub>): 6 0.8-1.0 (m, 2H), 1.0-1.9 (m,

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13C NMR (75 MHz, DMSO- $d_6$ ): amidine and carbonyl carbons: 6 165.8, 168.8, 169.9, 172.2 and 172.4.

MS m/z 500 (M+ 1)

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Example 32

Mecoc-CH2-CO-(R) Cha-Pic-Pab 20

See Example 31 (ii) above.

Example 33

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H2N-CO-CH2-(R) Cha-Pic-Pab

(i) H<sub>2</sub>N-CO-CH<sub>2</sub>-(R) Cha-Pic-Pab(Z)

40°C turned out to be an extremly sluggish reaction. Even the addition of 230 mg (2.6 mmol) lithium bromide did not seem to improve the reaction rate. However, addition of chloroacetamide in 3 ml acetonitrile in the presence of 395 mg (2.86 mmol) potassium carbonate by sonication at lithium iodide and heating/sonication gave small amounts Attempted alkylation of 455 mg (0.83 mmol) H-(R)Cha-Pic-Pab(Z) (See Example 25) with 80 mg (0.86 mmol) 30 35

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water, extraction with ethyl acetate/toluene, drying of the organic phase  $(MgSO_4)$  and removal of the solvent in vacuo gave a residue which was subjected to flash chromatography using MeOH/CH2Cl2 as eluent to give 118 mg of product, according to TLC. Workup by addition of (24%) of the desired product.

(ii)  $H_2N-CO-CH_2-(R)Cha-Pic-Pab \times 2 HCl$ 

143 mg 10% Pd/c for 2 h. The mixture was diluted with distilled water and hydrochloric acid and filtered 118 mg (0.2 mmol)  $\mathrm{H_2N-CO-CH_2-(R)\,Cha-Pic-Pab(Z)}$  dissolved through hyflo. Freeze drying gave 26 mg (24%) of the in 10 ml 95% ethanol was hydrogenated in the presence of desired product. 12 10

16H), 2.3 (d, 1H), 3.4 (t, 1H), 3.6 (AB-system, 2H), 3.8 H-NMR (300 MHz,  $CD_3OD$ ): 6 0.9-1.1 (m, 2H), 1.1-1.9 (m, (d, 2H), 4.35 (t, 1H), 4.5 (s, 2H), 5.2 (s, 1H), 7.55 (d,

2H), 7.8 (d, 2H).

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Example 34

Boc-(R) Cha-Pic-Pab

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presence of 38 mg 10% Pd/C for 4 h. Removal of the catalyst by filtration and evaporation of the solvent in vacuo followed by dissolution of the residue in water and 10 mg (0.015 mmol) Boc-(R)Cha-Pic-Pab(Z) (See Example 25) dissolved in 5 ml ethanol was hydrogenated in the freeze drying yielded 7.6 mg (95%) of the product.

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16H), 2.4 (d, 1H), 3.25 (t, 1H), 4.0 (d, 1H), 4.5 (AB- $^{1}_{H-NMR}$  (300 MHz,  $^{\circ}_{D_3}$ OD): 6 0.9-1.1 (m, 2H), 1.1-1.9 (m, system, 2H), 4.5-4.6 (m, 1H), 5.25 (s, 1H), 7.45 (d, 2H), 35

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#### Example 35

## Ac-(R)Cha-Pic-Pab x HCl

(i) Ac-(R)Cha-Pic-Pab(Z)

99.2/0.8 and 98.4/1.6) gave 0.24 g (60%) of the product. gradient of CH<sub>2</sub>Cl<sub>2</sub>/ MeOH ( 99.9/0.1, 99.8/0.2, 99.6/0.4, Evaporation and flash chromatography using a stepwise residue was dissolved in  $\mathrm{CH_2Cl_2}$  and washed with water. at room temperature the solvent was removed in vacuo. The temperature. After stirring for an additional 30 minutes 25) and 0.19 g (1.35 mmol)  ${
m K}_2{
m CO}_3$  in 10 ml  ${
m CH}_3{
m CN}$  at room of 0.37 g (0.68 mmol) H-(R)Cha-Pic-Pab(Z) (See Example Acetyl chloride 0.06 g (0.8 mmol) was added to a mixture

## (ii) Ac-(R)Cha-Pic-Pab x HCl

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title compound.  $\mathrm{NH_4OAC}$  followed by freeze drying from 1M HCl gave the (35/65) as eluent. Removal of the solvent and excess subjected to purification by RPIC using CH3CN/0.1 M NH4OAc and evaporation of the solvent the crude material was atmospheric pressure. After filtration of the catalyst Ac-(R) Cha-Pic-Pab(Z) was hydrogenated over 5 % Pd/C at

protons is totally obscured by the solvent-signal. 2H), 7.76 (d, 2H), 8.23 (m, 1H). The signal of one of the 1H), 4.46,4.57 (ABX, 2H), 5.16-5.22 (m, 1H), 7.51 (d, 19Н), 2.35-2.47 (ш, 1Н), 3.2-3.33 (ш, 1Н), 3.95-4.05 (ш, <sup>1</sup>H-NMR (300 MHz, CD<sub>3</sub>OD): & 0.85-1.1 (m, 2H), 1.15-2.0 É

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168.3, 172.5, 173.8, 175.1  $^{13}\mathrm{C-NMR}$  (75 MHz, CD $_3\mathrm{OD}$ ): amidine and carbonyl carbons:  $\delta$ 

MS m/z 456 (M++1)

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<u>Example 36</u>

## Me-802-(R) Cha-Pic-Pab x EC1

## (1) Me-SO<sub>2</sub>-(R)Cha-Pic-Pab(Z)

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the solvent in vacuo gave a residue which was subjected to flash chromatography using ethyl acetate/methanol with water followed by drying ( $Na_2SO_4$ ) and evaporation of (95/5) as eluent to give 159 mg (67%) of the product. was allowed to reach room temperature over night. Washing amine in 5 ml of methylene chloride. The reaction mixture Pab(Z) (See Example 25) and 0.11 ml (0.763 mmol) triethyl stirred solution of 209 mg (0.382 mmol) H-(R)Cha-Picin 0.5 ml methylene chloride was added at 0°C to a A solution of 48 mg (0.42 mmol) methanesulfonyl chloride

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## (ii) Me-SO<sub>2</sub>-(R)Cha-Pic-Pab x HCl

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25 20 mg (86%) of the product. was dissolved in 2 ml water and freeze dryed to give 116 evaporation of the solvent in vacuo gave a residue which 5 ml 95% ethanol and 1 ml water was hydrogenated in the 150 mg (0.24 mmol) Me-80 $_2$ -(R)Cha-Pic-Pab(z) dissolved in filtration, addition of 0.2 ml 1M hydrochloric acid and presence of 5% Pd/C for 4 h. Removal of the catalyst by

မ 3.35 (dt, 1H), 3.90 (bd, 1H), 4.45 (AB-system, 2H) 4.50-4.55 (m, 1H), 5.13 (dd, 1H), 7.50 (d, 2H), 7.75 (d, 2H). (m, 15H), 1.90 (bd, 1H), 2.30 (bd, 1H), 2.85 (s, 3H), <sup>1</sup>H-NMR (500 MHz, CD<sub>3</sub>OD): & 0.90-1.10 (m, 2H), 1.15-1.85

 $^{13}\mathrm{C\text{-}NMR}$  (125 MHz  $^{12}\mathrm{O}_2\mathrm{O}_3$ : amidine and carbonyl carbons:  $\delta$ 166.8, 173.0 and 174.6.

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#### Example 37

## H-(R)Cha-(R,8)betaPic-Pab x 2 HCl

5 (i) Boc-(R)Cha-(R,S)betaPic-Pab(Z)

EDC was added at -18°C to a stirred solution of 1.0 g (2.6 mmol) Boc-(R)Cha-(R,S)betaPic-OH (See preparation of starting materials), 1.28 g (10.5 mmol) DMAP, 0.74 g (2.6 starting materials), 1.28 g (10.5 mmol) DMAP, 0.74 g (2.6 mmol) H-Pab-(Z) (See preparation of starting materials) in 5 ml DMF. The reaction mixture was allowed to reach room temperature over night and the solvent was subsequently removed in vacuo. The residue was dissolved in CH<sub>2</sub>Cl<sub>2</sub> and removed in vacuo. The residue was dissolved in CH<sub>2</sub>Cl<sub>2</sub> and the organic layer was washed succesively with 0.3M KHSO<sub>4</sub>, the solvent gave a residue which was subjected to flash chromatography using heptane:ethyl acetate with 4% methanol as eluent to yield 0.74 g (44%) of the desired product.

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## (11) H-(R)Cha-(R,S)betaPic-Pab(Z)

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0.68 g (1.05 mmol) Boc-(R)Cha-(R,S)betaPic-Pab(Z) was dissolved in ethyl acetate saturated with HCl(g). The solution was stirred for 1 h at room temperature. Water was added and the mixture was made alkaline with K<sub>2</sub>CO<sub>3</sub>. The water phase was extracted with ethyl acetate. The organic phase was then washed with water and dried (Na<sub>2</sub>SO<sub>4</sub>). Evaporation gave 0.5 g (87%) of the desired product.

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## (iii) H-(R)Cha-(R,S)betaPic-Pab x 2 HCl

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65 mg (0.19 mmol) H-(R)Cha-betapic(R,S)-Pab(2) was dissolved in 7 ml ethanol and hydrogenated in presence of 5% Pd/C for 4 hours. Removal of the catalyst by filtration, evaporation of the solvent and freeze drying

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from 1M HCl and water gave 41 mg (71%) of the product.

<sup>1</sup>H NMR (300 MHz, D<sub>2</sub>O, 2 diastereomers 4/5, and rotamers); 6 0.8-2.16 (m, ), 2.5-2.77 (m, 3H), 3.13-3.43 (m, 3H), 5 3.68-3.94 (m, 1H), 4.18-4.41 (m, 1H), 4.41-4.52 (m, 3H), 7.46-7.57 (m, 2H), 7.72-7.83 (m, 2H).

#### Example 38

- 10 HOOC-CH2-CH2-(R)Cha-(R, S) betaPic-Pab x 2 HCl
- (i) Bnooc-CH2-CH2-(R)Cha-(R,S)betaPic-Pab(Z)

0.21 g (0.38 mmol) H-(R)Cha-(R,S)betaPic-Pab(2) (See Example 37) was dissolved in 2 ml ethanol. 0.68 g (0.42 mmol) benzyl acrylate was added and the solution was stirred for 5 days. Evaporation and flash chromatography with CH<sub>2</sub>Cl<sub>2</sub> /MeOH (95/5) as eluent gave 0.19 g (70%) of the desired product.

(ii) HOOC-CH2-CH2-(R)Cha-(R,S)betaPic-Pab x 2 HCl

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170 mg (0.24 mmol) BhOOC-CH<sub>2</sub>-CR<sub>2</sub>-(R)Cha-(R,S)betaPic-Pab(Z) was dissolved in 10 ml ethanol and hydrogenated in 25 presence of 5% Pd/C for 4 hours. Removal of the catalyst by filtration, evaporation of the solvent and freeze drying from 1M HCl and water gave 103 mg (77%) of the product.

- 30 <sup>1</sup>H NMTR (300 MHz, D<sub>2</sub>O,mixture of 2 diastereomers 4/5 and rotamers); 6 0.92-2.03 (m, H), 2.51-2.78 (m, 1H), 3.21-3.52 (m, 1H), 3.88-4.01 (m, 1H), 4.07-4.3 (m, 2H), 4.44-4.71 (m, 2H), 7.59 (d, 2H), 7.86 (d, 2H)
- 35 13C NWR (300.13 MHz, D<sub>2</sub>O, mixture of 2 diastereomers 4/5 and rotamers): amidine and carbonyl carbons: 6 167.0, 168.0, 168.1, 175.9, 176.0, 176.3, 176.4 and 178.2.

#### Example 39

### HOOC-CH2-(R)Cha-Val-Pab x 2 HCl

### (i) Boc-(R)Cha-Val-Pab(Z)

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gave 2.77 g (47%) of the desired product. combined organic extracts, removal of the solvent in vacuo and flash chromatography using  $ext{CH}_2 ext{Cl}_2/ ext{MeOH}$  as eluent and ethyl acetate. Subsequent drying (MgSO $_4$ ) of the with water was followed by extraction with toluene, ether reach room temperature over night and workup by dilution DMAP in 50 ml DMF. The reaction mixture was allowed to 3.41 g (9.2 mmol) Boc-(R)Cha-Val-OH(See preparation of preparation of starting materials), and 4.5 g (36.8 mmol) 1.77 g (9.2 mmol) EDC was added at -12°C to a mixture of starting materials), 2.61 g (9.2 mmol) H-Pab(Z) (See

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#### (ii) H-(R)Cha-Val-Pab(Z)

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ethyl acetate. Drying (potassium carbonate) and removal of the solvent in vacuo gave 1.8 g (77%) of H-(R) Cha-Valadded to pH 10 and the aqueous phase was extracted with acetate. After 15 minutes sodium carbonate solution was 9 (4.4 mmol) Boc- $\{R\}$ Cha-Val-Pab $\{Z\}$  in 75 ml ethyl Hydrogen chloride was bubbled through a solution of 2.77

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## (111) Bnooc-CH2-(R)Cha-Val-Pab(Z)

residue was purified by flash chromatography using filtered and the solvent was removed in vacuo. The in order to dissolve the product, and the mixture was sonicated for 2.5 h at 40°C. More acetonitrile was added, mmol) potassium carbonate in 2 ml acetonitrile was ml (0.67 mmol) benzyl bromoacetate, and 252 mg (1.83 A mixture of 326 mg (0.61 mmol) H-(R) Cha-Val-Pab(2), 105

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finally crystallised from ethyl acetate to give 124 mg methanol/methylene chloride as eluent. The product was (30%) of colourless crystals.

## (iv) HOOC-CH<sub>2</sub>-(R)Cha-Val-Pab x 2 HCl

15 10 mg (50%) of the desired compound. vacuo. Freeze drying of the remaining solution yielded 55 solvents were removed from the combined filtrates in was washed with dilute hydrochloric acid. The organic The mixture was filtered through hyflo and the filtercake hydrogenation was continued for another 2 hours at 50°C. 25 mg 10% Pd/C. 10 ml of THF was added and the ethanol was hydrogenated for 2 hours in the presence of 124 mg (0.18 mmol)  $BnOOC-CH_2-(R)Cha-Val-Pab(Z)$  in 20 ml

2H), 4.5 (m, 2H), 7.5 (s, 2H), 7.7 (s, 2H), 8.9 (s, 1H). 7H), 2.0-2.15 (bs, 1H), 3.45 (AB-system, 2H), 4.1 (m, <sup>1</sup>H-NMR (500 MHz, D<sub>2</sub>0); 6 0.75-1.4 (m, 12H), 1.5-1.9 (m,

#### Example 40

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## HOOC-CH2-CH2-(R) Cha-Val-Pab x 2 HC1

#### 25 (i) H-(R)Cha-(R,S)Val-Pab(Z)

30 title compound. protecting group was removed in the same way as described occured to give Boc-(R)Cha-(R,S)Val-Pab(Z). The Boc starting materials). A total epimerization of the valine described for Boc-(R)Cha-Pic-OMe (See preparation of Val-OH with H-Pab(Z), using the pivaloyl coupling as for Boc-(R)Cha-Val-Pab(Z) (See Example 39) to give the The title compound was prepared by coupling Boc-(R)Cha-

## (ii) BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R) Cha-(R,S) Val-Pab(Z)

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was kept at 40°C over night. The solvent was removed in and 308 mg (1.9 mmol) benzyl acrylate in 3 ml of ethanol the residue was purified by flash chromatography using methanol/methylene chloride (10/90) A solution of 1.007 g (1.9 mmol) H-(R)Cha-(R,S)Val-Pab(Z) as eluent to give 1.086 g (82%) of the title compound. vacuo and

(111) HOOC-CH2-CH2-(R)Cha-Val-Pab x 2 HCl

eluent. Two main fractions were isolated, of which the second fraction contained the title compound. 67 mg of celite and removal of the THF in vacuo followed by freeze drying of the remaining aqueous solution gave a residue of which approximately 300 mg was subjected to  $\mathtt{HPLC}$  using 25% acetonitrile in 0.1 M Ammonium acetate buffer as was hydrogenated in 25 ml THF and 14 ml 0.5 M hydrochloric acid in the presence of 223 mg 10% Pd/C for 2 hours. Removal of the catalyst by filtration through 1.086 g (1.6 mmol) BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cha-(R,S)Val-Pab(Z) title compound, as the dihydrochloride, was isolated. 20 13 2

7H), 1.65-1.9 (m, 7H), 2.15-2.25 (m, 1H), 2.85 (t, 2H), 3.15-3.2 (m, 1H), 3.3-3.35 (m, 1H), 4.15-4.2 (m, 1H),  $^{1}$ H-NWR (500 MHz, D<sub>2</sub>O); 6 1.0-1.15 (m, 12H), 1.2-1.4 (m, 4.25 (d, 1H), 4.55-4.65 (AB-system, 2H), 7.65 (d, 2H), 7.85 (d, 2H).

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 $^{13}\mathrm{C-NMR}$  (75 MHz,  $^{}\mathrm{D_2O}$ ): amidine and carbonyl carbons:  $^{\delta}$ 167.0, 169.8, 173.96 and 174.04.

Example 41

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H-(R)Hoc-Ase-Pab x 2 HCl

(i) Boc-(R)Hoc-Aze-Pab(Z) 32

Prepared in the same way as described for Boc-(R)Cha-Pic-

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with Boc-(R)Hoc-Aze-OH (See preparation of starting materials). The crude product was subjected to flash Pab(Z) (See Example 25) by replacing Boc-(R)Cha-Pic-OH chromatography (Toluene/EtOAc 1/6) to give 0.32 g (37%) of the desired product.

(ii) H-(R)Hoc-Aze-Pab(Z)

S

Boc-(R)Hoc-Aze-Pab(Z) was treated in the same way as described for Boc-(R)Cha-Pic-Pab(2) in Example 25 to to give 0.23 g (88%) of the title compound. 10

(iii) H-(R)Hoc-Aze-Pab x 2 HCl

in 3 ml ethanol and hydrogenated in presence of 5% Pd/C 20 mg (0.037 mmol) of H-(R)Hoc-Aze-Pab(2) was dissolved catalyst by filtration, evaporation of the solvent and for 4 hours at athmospheric pressure. Removal of the freeze drying from 1M HCl gave 11 mg (63%) of the 15 20

product.

 $^{1}\mathrm{H}$  NMR (300.13 MHz,  $\mathrm{D_{2}O},$  mixture of two rotamers 3:1): 4.65 (s, 2H), 5.0-5.11 (m, 1H), 7.62 (d, 2H), 7.9 (d, 2.7-3.0 (m, 1H), 4.1-4.3 (m, 1H), 4.35-4.56 (m, 1H), major rotamer: 6 0.9-2.1 (m, 15H), 2.4-2.6 (m, 1H), 2H). The signal of one of the protons is totally obscured by the H-O-D-signal.

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Example 42

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HOOC-CH2-CH2-(R) HOC-ARS-PAD X 2 TFA

(i) Bn00C-CH2-CH2-(R)Hoc-Aze-Pab(Z)

solution of 0.2 g (0.37 mmol) H-(R)Hoc-Aze-Pab(Z) (See Example 41) in 2 ml ethanol (95%) at room temperature. 0.067 g (0.41 mmol) benzylacrylate was added to a 35

to give 0.16 g (62%) of the desired product. purified with flash chromatography ( $CH_2Cl_2$ : MeOH, 96/4) The solvent was removed in vacuo and the residue was The reaction was left at room temperature for 5 days.

(ii) HOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Hoc-Aze-Pab x 2 TFA

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hydrogenation at atmospheric pressure in presence of 5% from water and TFA gave 120 mg (87%) of the product. filtration evaporation of the solvent and freeze drying Pd on charcoal for 3 hours. Removal of the catalyst by dissolved in 10 ml ethanol and subjected to 160 mg (0.23 mmol) BnOOC-CH $_2$ -CH $_2$ -(R)Hoc- $_2$ -Pab(Z) was

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4.35-4.58 (m, 2H), 4.65 (s, 2H), 5.0-5.12 (m, 1H), 7.63 б 0.9-1.9 (т, 13Н), 1.94-2.16 (т, 2Н), 2.38-2.55 (т,  $^{
m 1}{
m H}$  NMR (300.13 MHz, D $_{
m 2}{
m O}$  2 rotamers 3:1); major rotamer: (d, 2H), 7.87 (d, 2H) 1H), 2.7-2.97 (m, 3H), 3.2-3.44 (m, 2H), 4.16 (m, 1H),

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δ 167.3, 168.7, 172.5 and 176.6.  $^{13}\mathrm{C}$  NMR (300.13 MHz, D<sub>2</sub>0): amidine and carbonyl carbons: 20

Example 43

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HOOC-CH2-(R, 8) CH(COOH)-(R) Hoc-Pro-Pab x 2 HC1

(i) Boc-(R)Hoc-Pro-Pab(Z)

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title compound. using ethyl acetate as eluent gave 0.886 g (58 %) of the starting materials) in the same way as described for Boc-Prepared from Boc-(R)Hoc-Pro-OH (See preparation of (R)Cha-Pic-Pab(Z) in Example 25. Flash chromatography

27H (thereof 1.2 (s, 9H)), 2.1-2.4 (m, 1H), 3.3-3.5 (m, 1H-NMR. (300 MHz, CDCl<sub>3</sub>); & 0.7-0.95 (m, 2H), 0.95-2.1 (m, **3**5

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9.4 (bs, 1H). (d, 1H), 7.1-7.4 (m, 7H), 7.65 (m, 1H), 7.7-7.8 (d, 2H), 2H), 4.45-4.6 (d, 1H), 5.15 (apparent bs, 2H), 5.2-5.3 1H), 3.65-3.95 (m, 1H), 4.0-4.2 (m, 1H), 4.2-4.45 (m,

 $^{13}\mathrm{C-NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $\ell$ 156.3, 164.6, 168.1, 171.4 and 172.4.

10 (ii) H-(R)Hoc-Pro-Pab(Z)

20 15 compound in almost quantitative yield. organic phase was washed with brine and dried ( $Na_2SO_4$ ) and the solvent evaporated in vacuo to give the title carbonate was added and the phases were separated. The evaporated and ethyl acetate and saturated sodium reaction mixture during 5 minutes. The solvent was and therefore hydrogen chloride was bubbled through the temperature. The reaction was not completed after 1.5 h 0°C. The temperature was allowed to rise to roomadded to 0.82 g (1.266 mmol) Boc-(R)Hoc-Pro-Pab(Z) at 40 ml ethyl acetate saturated with hydrogen chloride was

25 7.35-7.45 (m, 2H), 7.6-7.7 (m, 1H), 7.7-7.85 (d, 2H). 2H), 4.5-4.6 (m, 1H), 5.15 (s, 2H), 7.15-7.35 (m, 5H), 17H), 3.3-3.55 (m, 2H), 3.55-3.7 (m, 1H), 4.25-4.45 (m, <sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>); 6 0.75-0.95 (m, ZH), 0.95-2.4 (m,

30  $^{13}\mathrm{C-NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $_6$ 164.5, 167.8, 171.4 and 175.3.

(iii) BnOOC-CH<sub>2</sub>-(R,S)CH(COOBn)-(R)HoG-Pro-Pab(Z)

**3**5 To 0.15 g (0.5 mmol) benzyl acrylate in 1.5 ml EtOH (99%) the mixture was stirred at room temperature for 10 days. was added 0.273 g (0.498 mmol) H-(R)Hoc-Pro-Pab(Z) and

The solvent was removed in vacuo and the residue was subjected to flash chromatography, using ethyl acetate as eluent to give 0.103 g (25 %) of BnooC-CH<sub>2</sub>-(R,S)CH(COOBn)-

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(R)Cha-Pic-Pab(Z) (See Example 25). Flash chromatography starting materials) and H-Pab(Z) (See preparation of starting materials) in the same way as described for Bocusing ethyl acetate as eluent gave 1.3 g (78 %) of the 2

title compound.

1H-NMR (300 MHz, CDCl<sub>3</sub>): 6 0.75-0.95 (m, 2H), 0.95-2.0 (m, 7.15-7.5 (m, 7H), 7.7-7.85 (d, 2H), 9.45 (bs, 1H).

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 $^{13}\mathrm{C-NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $^6$ 156.6, 164.7, 168.1, 170.0 and 173.0.

(ii) H-(R)Hoc-Pic-Pab(Z)

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brine and dried  $(\mathrm{Na}_2\mathrm{SO}_4)$  and the solvent evaporated in acetate and saturated sodium carbonate was added and the phases were separated. The organic phase was washed with 100 ml ethyl acetate saturated with hydrogen chloride was The solvent was evaporated after 40 minutes and ethyl added to 1.3 g (1.96 mmol) Boc-(R)Hoc-Pic-Pab(Z) at 0°C. The temperature was allowed to rise to room-temperature. vacuo to give 0.85 g (77.5 %) of the product.

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2H), 5.15 (apparent bs, 3H), 7.05-7.2 (d, 2H), 7.2-7.35 1H-NMR (300 MHz, CDCl3): 6 0.75-0.95 (m, 2H), 1.05-2.3 (m, 25H), 3.0-3.15 (m, 1H), 3.6-3.75 (m, 2H), 4.25-4.4 (m, (m, 4H), 7.35-7.4 (d, 1H), 7.6-7.8 (d, 2H). 35

4.9-5.1 (m, 3H), 5.2 (s, 2H), 7.1-7.2 (m, 1H), 7.2-7.4 (m, 13H), 7.4-7.45 (d, 2H), 7.6-7.8 (m, 3H).

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<sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>); 6 0.75-2.05 (m, 18H), 2.3-2.45 (m, 1H), 2.45-2.8 (m, 3H), 3.15-3.45 (m, 3H), 3.5-3.65

(R)Hoc-Pro-Pab(Z).

(m, 1H), 4.3-4.5 (m, 2H), 4.55-4.7 (m, 1H), 4.8 (s, 1H),

(iv)  $HOOC-CH_2-(R,S)CH(COOH)-(R)HOC-Pro-Pab x 2 HCI$ 

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% Pd/C for 2 h. Removal of the catalyst by filtration Pab(Z) dissolved in 4 ml ethanol (99.5 %) and 0.3 ml chloroform was hydrogenated in the presence of 111 mg 5 and evaporation of the solvent followed by dissolving in water and freeze drying showed incomplete hydrogenation. The hydrogenation was continued in the presence of 103 mg (0.122 mmol) BnOOC-CH2-(R,S)CH(COOBn)-(R)Hoc-Proethanol, 1 N HCl and 5 % Pd/C for 5 hours. 20

Removal of the catalyst by filtration and evaporation of the solvent followed by dissolving in water and freeze drying gave the title compound.  $^{1}\mathrm{H-NMR}$  (500 MHz, CD30D, mixture of two diastereomers);  $\delta$ 2.15 (m, 5H) 2.25-2.35 (m, 1H), 2.9-3.2 (m, 2H), 3.5-3.65 0.8-1.0 (m, 2H), 1.1-1.4 (m, 6H), 1.6-1.8 (m, 5H), 1.9-(m, 1H), 3.7-3.9 (2m, total 1H), 4.15-4.4 (2m, total 1H), 4.4-4.6 (п, 4Н), 7.5-7.6 (п, 2Н), 7.7-7.85 (п, 2Н). 13C-NWR (75 MHz, CDCl3): amidine and carbonyl carbons: 6 167.9, 168.2, 168.3, 172.8, 173.6, 174.3 and 174.4.The the two diastereomers are partly 35

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Example 44

(i) Boc-(R)Hoc-Pic-Pab(Z) S

Prepared from Boc-(R)Hoc-Pic-OH (See preparation

1H), 3.8 (m, 1H), 4.2-4.45 (m, 2H), 4.45-4.55 (m, 2H), 5.15 (apparent bs, 3H), 5.25-5.3 (m, 1H), 7.0 (bs, 1H), 31H (thereof 1.3 (s, 9H)), 2.4-2.5 (m, 1H), 3.0-3.1 (m,

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164.5, 167.9, 170.8 and 175.7.  $^{13}\mathrm{C-NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $\delta$ 

## (iii) BnOOC-CH2-(R)Hoc-Pic-Pab(Z)

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second compound eluated was the title compund (0.27 g). column was  $(BnOOC-CH_2)_2(R)Hoc-Pic-Pab(Z)$  (0.28 g) and the give 2 products. The first compound eluated from the flash chromatography using ethyl acetate as eluent, to vacuo gave 0.626 g of a residue which was subjected to was washed with brine and dried ( $Na_2SO_4$ ). Evaporation in added. The phases were separated and the organic phase solvent was removed and ethyl acetate and water was mixture was heated to 60°C in oilbath for 1 h. 0.217 g (1.57 mmol)  $K_2CO_3$  in 7 ml acetonitrile. mixture of 0.4 g (0.712 mmol) H-(R)Hoc-Pic-Pab(Z) and 0.171 g (0.748 mmol) benzyl bromoacetate was added to a The

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20  $BnOOC-CH_2-(R)Hoc-Pic-Pab(Z)$ :

5.3 (m, 1H), 7.1-7.45 (m, 12H), 7.7-7.8 (d, 2H). system, 2H), 4.75 (s, 2H), 5.15 (apperent s, 3H), 5.25-1H), 3.35-3.5 (m, 2H), 3.6-3.7 (m, 1H), 4.35,4.55 (ABX-18H), 2.3-2.5 (m, 1 or 2H), 2.9-3.05 (m, 1H), 3.2-3.3 (m,  $^{1}\mathrm{H\text{-}NMR}$  (300 MHz, CDCl $_{3}$ ): & 0.7-0.95 (m, 2H), 1.0-1.75 (m,

25

164.6, 167.9, 170.5, 173.4 and 175.0.  $^{13}\mathrm{C\text{-}NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $\epsilon$ 

(iv) HOOC-CH<sub>2</sub>-(R)Hoc-Pic-Pab x 2 HC]

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for 4 h. Removal of the catalyst by filtration and evaporation of the solvent followed by dissolving in (1 N) was hydrogenated in the presence of 280 mg 5 % Pd/C in 7.8 ml ethanol (99.5 %) and 1.2 ml hydrogen chloride 259 mg (0.365 mmol)  $BnOOC-CH_2-(R)Hoc-Pic-Pab(Z)$  dissolved

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The state of the same

water and freeze drying gave 170 mg (83 %) of the title

167.8, 168.6, 169.6 and 172.3.  $^{13}\mathrm{C-NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $\delta$ 4.3-5.05 (m, 2H), 7.1-7.4 (m, 2H), 7.4-7.7 (m, 2H). 1H), 2.9-3.2 (m, 1H), 3.4-3.9 (m, 3H), 4.05-4.3 (m, 2H), <sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>): & 0.4-1.85 (m, 20H), 1.85-2.2 (m,

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#### Example 45

## (HOOC-CH2)2-(R)Hoc-Pic-Pab x 2 HC1

## (i) (BnOOC-CH<sub>2</sub>)<sub>2</sub>(R)Hoc-Pic-Pab(Z)

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The title compound was obtained in the alkylation of H-(R)Hoc-Pic-Pab(Z) as described in Example 44 above.

25 20 5.35 (m, 1H), 7.1-7.45 (m, 16H), 7.5-7.65 (m, 1H), 7.7-7.85 (d, 2H). 6H), 4.35-4.55 (m, 2H), 4.9 (2s, 4H), 5.2 (s, 2H), 5.25-18H), 2.35-2.5 (m, 1H), 2.9-3.05 (m, 1H), 3.5-3.85 (m, <sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>): & 0.7-0.95 (m, 2H), 0.95-1.95 (m,

164.7, 167.9, 170.5, 172.0 and 172.4.  $^{13}\mathrm{C-NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $\delta$ 

## (ii) (HOOC-CH<sub>2</sub>)<sub>2</sub>-(R)Hoc-Pic-Pab x 2 HCl

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 $^{
m CH_2})_2-({
m R})$  Hoc-Pic-Pab dihydrochloride. This crude material and evaporation of the solvent followed by dissolving in water and freeze drying gave 109 mg (99 %) of (HOOC-5 % Pd/C for 3.5 h. Removal of the catalyst by filtration chloride (1 N) was hydrogenated in the presence of 150  $_{
m mg}$ dissolved in 4.5 ml ethanol (99.5 %) and 0.5 ml hydrogen 153 mg (0.178 mmol) (BnOOC-CH<sub>2</sub>)<sub>2</sub>-(R)Hoc-Pic-Pab(Z)

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(80 % purity) was subjected to putification by RPLC using CH3CN/0.1 M NH40Ac, 1:4 as eluent. Removal of the solvent and excess NH40Ac followed by freeze drying from 1 M HCl gave the title compound.

rotamer: 6 0.95-2.15 (m, 20H), 2.25-2.35 (m, 1H), 3.45-3.55 (т, 1Н), 3.95-4.25 (т, 5Н), 4.6-4.65 (т, 2Н), 4.92-5.01 (m, 1H) 5.15-5.20 (m, 1H), 7.58-7.63 (d, 2H), 7.84- $^{\mathrm{H-NMR}}$  (500 MHz,  $^{\mathrm{D}_2\mathrm{O}}$ , mixture of two rotamers): 7,89. (d, 2H).

Resolved signals arising from the minor rotamer appears at: 6 0.7-0.85 (m), 2.35-3.45 (m), 3.05-3.15 (m), 4.47-4.55 (m), 4.55-4.6 (m), 4.65-4.7 (m), 7.63-7.67 (d), 7.89-7.95 (d).

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 $^{13}\mathrm{C-NMR}$  (75 MHz,  $^{12}\mathrm{O}_2\mathrm{O}$ ): amidine and carbonyl carbons:  $^6$ 168.20, 169.70, 170.20 and 172.71.

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Example 46

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HOOC-CH2-(R) Pro(3-(S) Ph) -Pro-Pab x 2 HCl

(i) Boc-(R)Pro(3-(S)Ph)-Pro-Pab(Z)

25

1 x 20 + 1 x 10 ml 1M  $\mathrm{XHSO}_4$ , 1 x 15 ml  $\mathrm{NaHCO}_3(\mathrm{aq})$ , 3 x 15 ml water, 1 x 15 ml brine and dried (MgSO $_4$ ). Filtration (1.5/1) was added 310 mg (1.62 mmol) EDC and the mixture was stirred for 23 h at room temperature. Most of the solvent was evaporated and 50 ml water was added to the residue. The water phase was extracted with 1  $\times$  75 and 2  $\kappa$  50 ml EtOAc. The combined organic phase was washed with and evaporation of the solvent gave 670 mg of an oil materials) and 733 mg (6 mmol) DWAP in 25 ml  ${
m CH_3CN/DMF}$ Pro-OH (See preparation of starting materials), 425 mg (See preparation of starting To a solution of 570 mg (1.5 mmol) Boc-(R)Pro(3-(S)Ph)-(1.5 mmol) H-Pab(Z)

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which was purified by flash chromatography using EtOAc as eluent which gave 529 mg (55 %) of the title compound.

1H), 5.19 (8, 2H), 7.1-7.37 (m, 10H), 7.42 (d, 2H), 7.81 3H), 2.1-2.31 (m, 3H), 2.52 (q, 1H), 3.58-3.77 (m, 4H), <sup>1</sup>H-NWR (300 MHz, CDCl<sub>3</sub>): 6 1.26 (s, 9H), 1.53-1.88 (m, 4.31 (d, 1H), 4.35 and 4.47 (ABX-system, 2H), 4.65 (dd, (d, 2H), 8,0 (t, 1H (NH)).  $^{13}\mathrm{C-NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $^6$ 154.6, 164.6, 168.1, 171.1 and 171.3. 2

(11) H-(R) Pro(3-(S) Ph) -Pro-Pab(Z)

the organic phase was washed with 1  $\times$  10 ml 2 M NaOH, 1 Filtration and evaporation of the solvent gave 403 mg (90 temperature and stirred for 3 h. The solvent was 529 mg (0.81 mmol) of Boc-(R)Pro(3-(S)Ph)-Pro-Pab(Z) was dissolved in 15 ml EtOAc/HCl(g,saturated) at room evaporated and the residue was dissolved in 70 ml  ${
m CH_2CL_2}$ . x 10 ml water, 1 x 10 ml brine and dried (MgSO<sub>4</sub>). %) of the title compound as a white powder. 20 12

4H), 3.83 (bd, 1H), 4.25-4.45 (m, 2H), 4.53 (m, 1H), 5.19 <sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>): 6 1.44-1.57 (m, 1H), 1.62-1.86 (m, 2H), 1.96-2.35 (m, 3H), 2.45 (q, 1H), 3.05-3.35 (m, (s, 2H), 7.16-7.37 (m, 10H), 7.42 (d, 2H), 7.66 (t, 1H, (NH)), 7.77 (d, 2H). 25

 $^{13}\mathrm{C-NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $\delta$ 164.4, 167.9, 171.1 and 173.0. 8

(111) BnOOC-CH2-(R)Pro(3-(S)Ph)-Pro-Pab(Z)

Pab(Z), 105 mg (0.46 mmol) Br-CH2-COOBn and 125 mg (0.90 mmol) K<sub>2</sub>CO<sub>3</sub> in 10 ml CH<sub>3</sub>CN was heated to 50°C for 1 h and A mixture of 200 mg (0.36 mmol) H-(R)Pro(3-(S)Ph)-Pro-35

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a stepwise gradient of CH<sub>2</sub>Cl<sub>2</sub>/MeOH(NH<sub>3</sub>-saturated) (95/5 crude material was purified by flash chromatography using followed by 9/1) to give 182 mg (72 %) of the title compound as a white solid. evaporation of the solvent gave 260 mg of an oil. The washed with 10 ml water and dried (MgSO $_4$ ). Filtration and was dissolved in 70 ml EtOAc. The organic phase was 30 minutes. The solvent was evaporated and the residue

10 (m, 15H), 7.43 (d, 2H), 7.5-7.8 (m, 3H, one NH). system centered at 4.37, 2H), 4.58 (dd, 1H), 4.97-5.1 <sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>): & 1.43-1.82 (m, 3H), 1.96-2.13 (AB-system centered at 5.03, 2H), 5.19 (s, 2H), 7.16-7.38 (m, 2H), 3.24-3.51 (m, 4H), 3.83 (d, 1H), 4.29-4.46 (ABX-(m, 1H), 2.14-2.22 (m, 1H), 2.26-2.43 (m, 2H), 3.02-3.14

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 $^{13}\mathrm{C-NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $\delta$ 164.5, 167.9, 171.15, 171.2 and 172.7.

20 (iv) HOOC-CH<sub>2</sub>-(R)Pro(3-(S)Ph)-Pro-Pab x 2 HCl

NH4OAc/CH3CN 4/1 followed by 3/1. Evaporation followed by gave 129 mg of a crude product . The crude product was (50%) of the pure product. freeze drying from water and 1N HC1-solution gave 70 mg purified by RPLC using a stepwise gradient of 0.1 M the solvent followed by freeze drying twice from water Filtration of the catlyst through hyflo, evaporation of was hydrogenated at atmospheric pressure for one hour. solution, 1 ml water and 10 ml ethanol and the mixture Pab(Z) was mixed with 0.075 g 5 % Pd/C, 1.0 ml 1N HCl-0.18 g ( 0.26 mmole ) of BnOOC-CH<sub>2</sub>-(R)Pro(3-(S)Ph)-Pro-

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1H),, 1.83-1.98 (m, 1H), 2.03-2.20 (m, 2H), 2.63 (t, 2H), system central at  $\delta$  3.88, 2 H), 4.06-4.19 (m, 1H), 4.37-3.28-3.40 (m, 1H), 3.55-3.78 (m, 2H), 3.81-3.96 (AB-1H-NMR (300 MHz, D<sub>2</sub>0): & 1.42-1.60 (m, 1H), 1.65-1.83 (m,

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4.70 (d, 1H), 7.35-7.58 (m, 7H), 7.74 (d, 2H) 4.61 (AB-system central at & 4.49, 2 H), 4.48 (dd, 1H),

167.02, 167.2, 169.3 and 174.4.  $^{13}\mathrm{C\text{--}NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons:  $\delta$ 

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#### Example 47

HOOC-CH2-CH2-(R) Pro(3-(8) Ph) - Pro-Pab x 2 HC1

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(i) BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Pro(3-(S)Ph)-Pro-Pab(Z)

20 15 mg (83 %) of the title compound.  $\mathrm{CH_2Cl_2/MeOH(NH_3-saturated)}$  (95/5 followed by 9/1) gave 202 chromatography using Evaporation of mixture was stirred at room temperature for 24 h. 114 mg (0.70 mmol) of benzyl acrylate and the reaction Pro-Pab(Z) (See Example 46) in 7 ml EtOH (99%) was added To a solution of 190 mg (0.34 mmol) H-(R)Pro(3-(S)Ph)the solvent a stepwise followed by gradient flash

25 7.75~7.85 (m, 3H, one NH). 2H), 5.19 (s, 2H), 7.15-7.37 (m, 15H), 7.44 (d, 2H), 2H), 4.61 (m, 1H), 4.48-5.08 (AB-system centered at 5.03, 2H), 2.84-2.96 (m, 1H), 3.18-3.48 (m, 4H), 4.28-4.44 (m, 1H), 1.9-2.05 (m, 1H), 2.2-2.64 (m, 5H), 2.69-2.82 (m, 1H-NMR (300 MHz, CDCl<sub>3</sub>): & 1.5-1.71 (m, 2H), 1.74-1.9 (m,

30 164.6, 168.0, 171.2, 172.5 and 172.9.  $^{13}\mathrm{C\text{-}NMR}$  (75 MHz,  $\mathrm{CDCl}_3$ ): amidine and carbonyl carbons:  $\delta$ 

(ii) HOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Pro(3-(S)Ph)-Pro-Pab x 2 HCl

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0.20 g ( 0.28 mmole ) of BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Pro(3-(S)Ph)-HCl-solution, 1 ml water and 10 ml ethanol and the Pro-Pab(Z) was mixed with 0.075 g 5 \$ Pd/C, 1.0 ml 1N

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evaporation of the solvent followed by freeze drying mixture was hydrogenated at atmospheric pressure for one hour. Filtration of the catalyst through hyflo, twice from water gave 125 mg 79 % of the title compound. H-NMR (300 MHz, D<sub>2</sub>0): 6 1.44 (m, 1H), 1.65-1.9 (m, 2H), 2.0-2.2 (m, 2H), 2.62 (q, 2H), 2.83 (t, 2H), 3.27-3.4 (m, 1H), 3.4-3.8 (m, 4H), 4.0-4.15 (m, 1H), 4.35-4.6 (m, 3H), 4.68 (d, 1H), 7.35-7.6 (m, 7H), 7.77 (d, 2H)

13C-NWR (75 MHz, CDCl3): amidine and carbonyl carbons: 166.2, 167.1, 174.1 and 174.2.

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#### Example 48

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HOOC-CH2-CH2-(R)Tic-Pro-Pab x 2 HCl

### (i) Boc-(R)Tic-Pro-Pab(Z)

Pab(2) (See Example 25) using Boc-(R)Tic-Pro-OH(See preparation of starting materials) instead of Boc-(R)Cha-Pic-OH. Flash chromatography using heptane/EtOAc (4/1) followed by EtOAc as eluents gave 425 mg (37%) of the Prepared in the same way as described for Boc-(R)Cha-Pictitle compound. 52 20

<sup>1</sup>H NWR (500 MHz, CDCl<sub>3</sub>): 6 1.35 (s, 9H), 1.95-2.15 (m, 3H), 2.4 (m, 1H), 2.8 (m, 1H), 3.3 (m, 1H), 3.55 (m, 2H), 4.25-4.4 (two m, 2H), 4.55-4.7 (two m, 2H), 7.15-7.5 (m, 10H), 7.85 (d,2H).

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6 164.6, 171.5 and 171.6. (two peaks are probably 13C-NWR (75.0 MHz, CDCl3): amidine and carbonyl carbons: overlapping)

(ii) H-(R)Tic-Pro-Pab(Z)

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in EtOAc saturated with HCl(g) and stirred at room Boc-(R)Tic-Pro-Pab(Z) (379 mg, 0.59 mmol) was dissolved temperature. Evaporation of the solvent gave 251 mg (79%) of the title compound as a white powder.

1H), 3.55 (m, 1H), 3.85 (m, 1H), 4.35-4.55 (m, 2H), 4.75 <sup>1</sup>H NMR (500 MHz, CDCl3): δ 1.65-2.15 (two m, 7H), 2.45 (m, 1H), 2.75 (m, 1H), 2.9 (m, 1H), 3.0 (m, 1H), 3.25 (m, (d, 1H), 4.9 (s, 1H), 5.25 (s, 2H), 6.8-7.45 (several m, 8H), 7.5 and 7.85 (two d, 4H).

6 164.5, 171.3 and 172.7 (two peaks are probably 13C-NMR (75.0 MHz, CDCl3): amidine and carbonyl carbons:

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overlapping).

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(iii) BnO<sub>2</sub>C-CH<sub>2</sub>-CH<sub>2</sub>-(R)Tic-Pro-Pab(Z)

20°C during 48 h. Evaporation of the solvent and flash chromatography using (50% EtOAc/Heptan then 10% MeOH/EtOAc) as eluent afforded 133 mg (73%) of the benzyl acrylat (63 mg, 0.39 mmol) in EtOH (1.3 ml) at H-(R)Tic-Pro-Pab(Z) (140 mg, 0.26 mmol) was treated with desired product as a white solid material.

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2H), 5.25 (s, 2H), 6.85-7.45 (several m, 12H), 7.5 and 2H), 3.9 (m, 1H), 4.45 (m, 2H), 4.65 (m, 1H), 5.1 (two d,  $^{1}\mathrm{H}$  NMR (500 MHz, CDCl $_{3}$ ): 6 1.75-2.0 (two m, 4H), 2.25 (m, 1H), 1.4-1.65 (m, 3H), 2.7-2.95 (two m, 4H), 3.05-3.2 (m, 7.9 (two d, 4H).

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6 171.5, 171.9 and 172.1 (two peaks are probably 13C-NMR (75.0 MHz, CDCl3): amidine and carbonyl carbons: overlapping).

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(iv) HOOC-CH2-CH2-(R)Tic-Pro-Pab x 2 HCl

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 $^{1}H$  NMR (500 MHz, D<sub>2</sub>O): & 2.1-2.35 (two m, 3H), 2.6 (m, 1H), 2.95-3.1 (m, 2H), 3.25-3.5 (two m, 2H), 3.65 (m, 3H), 4.65 (s, 2H), 4.75 (m, 1H), 5.85 (s, 1H), 7.15-7.6 (three m, 4H), 7.6 and 7.85 (two d, 4H).

 $^{13}\text{C-NMR}$  (75.0 MHz,  $D_2\text{O})$ : amidine and carbonyl carbons: & 166.9, 167.1 and 174.3 (two peaks are probably overlapping).

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#### Example 49

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## HOOC-CH2-CH2-(R)Cgl-Aze-Pig x 2 HCl

### (i) Boc-(R)Cgl-Aze-Pig(Z)2

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To a mixture of 0.623 g (1.83 mmole) Boc-(R)Cgl-Aze-OH(See preparation of starting materials), 0.816 g (1.92 mmole) H-Pig(Z)<sub>2</sub> (See preparation of starting materials) and 0.89 g (7.3 mmole) DMAP in 10 ml dichloromethane was added 0.368 g (1.92 mmole) of EDC and the mixture was stirred over night. The mixture was diluted and washed with 0.3 M KHSO<sub>4</sub> and once with brine. The organic layer was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and evaporated to yield 1.4 g of a crude product. Purification by flash chromatography using ethyl acetate as eluent gave 0.3 g (22%) of the pure product.

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### (ii) H-(R)Cgl-Aze-Pig(Z)2

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0.3 g (0.4 mmole) Boc-(R)Cgl-Aze-Pig(Z) $_2$  was mixed with 10 ml dichloromethane and 2.5 ml trifluoroacetic acid. The mixture was stirred for one and a half hour. After

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evaporation of the solvent the residue was dissolved in dichloromethane and washed twice with 0.2 M NaOH-solution. The combined water layer was extracted one more time with dichloromethane. The combined organic layer was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and evaporated to yield 0.24 g (93%) of the product.

1H-NMR (300 MHz, CDCl<sub>3</sub>, 339K): 6 0.9-1.9 (m, 15H), 1.94 (bd, 1H), 2.37-2.52 (m, 1H), 2.65-2.8 (m, 1H), 2.9-3.08 (m, 3H), 3.20 (t, 2H), 4.05-4.28 (m, 4H), 4.86 (dd, 1H), 5.16 (s, 4H), 7.2-7.42 (m, 10H), 7.98 (bs, NH).

## (iii) Bnooc- $CH_2$ - $CH_2$ -(R) $Cgl-Aze-Pig(Z)_2$

0.231 g (0.36 mmole) was dissolved in 2 ml ethanol and 61 μl (0.40 mmole) bensylacrylate was added. The reaction mixture was stirred for five days at room temperature. The mixture was evaporated and the crude product purified by flash chromatography using a stepwise gradient of CH<sub>2</sub>Cl<sub>2</sub>/MeOH (95/5, 90/10) as eluent to yield 0.218 g (75%) of the pure product.

1H-NMR (300 MHz, CDCl<sub>3</sub>, 335K): \$ 0.93 (bq, 1H), 1.02-1.85 (m, 14H), 1.94 (bd, 1H), 2.33-2.5 (m, 3H), 2.58-2.77 (m, 2H), 2.79-3.02 (m, 4H), 3.17 (t, 2H), 4.0-4.25 (m, 4H), 4.86 (dd, 1H), 5.11 (s, 2H), 5.12 (s, 4H), 7.2-7.4 (m, 15H), 8.03 (bs, NH), 10.35 (bs, NH)

# 30 (iv) $HOOC-CH_2-CH_2-(R)Cgl-Aze-Pig \times 2 HCl$

0.218 g ( 0.27 mmole ) of Bnooc-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cgl-Aze-Pig(Z)<sub>2</sub>
was mixed with 0.10 g 5 % Pd/C, 1 ml 1M HCl-solution, 1
ml water and 10 ml ethanol and the mixture was
hydrogenated at atmospheric pressure for one hour.
Filtration of the catalyst through hyflo, evaporation of
the solvent followed by freeze drying twice from water

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gave 134 mg (95%) of the title compound.

9H), 2.22-2.34 (m, 1H), 2.61-2.76 (m, 1H), 2.88 (t, 2H), 3.08 (bt, 2H), 3.19 (d, 2H), 3.34 (m, 2H), 3.83 (bd, 2H), <sup>1</sup>H-NMR (300 MHz, D<sub>2</sub>0): 6 1.0-1.4 (m, 7H), 1.55-2.05 (m, 3.95 (d, 1H), 4.29-4.49 (m, 2H), 4.90 (dd, 1H)  $^{12}\mathrm{C-NMR}$  (75 MHz,  $^{}\mathrm{D_2O}$ ): amidine and carbonyl carbons:  $^{\delta}$ 156.4, 167.6, 172.1 and 174.7

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Example 50

HOOC-CH2-(R)CG1-Pro-Pig x 2 HCl

(i) Boc-(R)Cgl-Pro-Pig(Z)2 12

using ethylacetate as eluent. This gave two products; 720 mg (34 %) of the title compound which eluted first from the column followed by 775 mg (44 %) of Boc-(R)Cgl-Prowas added. The phases were separated and the organic drying (Na $_2$ SO $_4$ ) and evaporation of the solvents gave 2.033 g of a residue wich was subjected to flash chromatography to rise to roomtemperature over night. The solvent was evaporated in vacuo and methylenchloride and 1 M XHSO4 starting materials), 1.197 g (2.82 mmol) H-Pig( $\mathbf{Z}$ )<sub>2</sub> (See preparation of starting materials) and 1.38 g (11.28 mmol) DMAP in acetonitrile. The temperature was allowed phase was washed with saturated  $NaHCO_3$ , water and brine, Pig(Z) formed by loss of one of the 2-protecting groups. 0.568 g (2.96 mmol) EDC was added at -15°C to a mixture of 1 g (2.82 mmol) Boc-(R)Cgl-Pro-OH (See preparation of

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ring, wich exhibit a broad peak ranging from 3.5 to 4.5 <sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>); Some signals, especially in the piperidin ring, are selectively broader due to an intramolecular exchange process. This is especially pronounced for the 2- and 6-CH $_{
m 2}$  groups of the piperidin

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ndd.

(m, 1H), 3.97-4.1 (m, 1H), 4.52-4.62 (d, 1H), 5.1 3.45-3.55 (m, 1H), 3.55-3.65 (m, minor rotamer), 3.8-3.93 6 0.85-2.1 (m, 19H), 2.3-2.45 (m, 1H), 2.8-3.2 (m, 4H), (apparent bs, 5H), 7.12-7.41 (m, 10H).

'n

13C-NMR (75 MHz, CDCl3): amidine and carbonyl carbons: 6 155.2, 156.3, 171.0 and 172.1.

(ii) H-(R)Cgl-Pro-Pig(Z)2

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ethylacetate and 2M NaOH was added. The organic layer was dissolved in 35 ml of  $TFA/CH_2Cl_2$ , 1/4 and stirred for 30 The solvent was removed in vacuo and washed with water and brine, dried  $(\mathrm{Na_2SO_4})$  and the solvent was evaporated in vacuo to give the title (0.946 mmol) of Boc-(R)Cgl-Pro-Pig(Z)2 was compound in quantitative yield. minutes. 720 mg

13

piperidin ring, are selectively broader due to an intramolecular exchange process. This is especially pronounced for the 2- and 6-CH $_2$  groups of the piperidin ring, wich exhibit a broad peak ranging from 3.5 to 4.5  $^{1}\mathrm{H-NMR}$  (300 MHz, CDCl $_{3}$ ); Some signals, especially in the

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2.98-3.18 (m, 2H), 3.18-3.35 (m, 1H), 3.35-3.5 (qvart., 6 0.8-2.15 (m, 19H), 2.22-2.4 (m, 1H), 2.75-2.98 (m, 2H), 1H), 3.5-3.7 (m, 1H), 4.42-4.58 (d, 1H), 5.1 (s, 4H), 7.1-7.5 (m, 10H).

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 $^{13}\mathrm{C-NMR}$  (75 MHz, CDCl $_3$ ): amidine and carbonyl carbons;  $\delta$ 154.96, 171.31, 174.82.

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(iii) Bnooc-CH<sub>2</sub>-(R)Cgl-Pro-Pig(Z)<sub>2</sub>

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the column and 142 mg (18 %) of the title compound. ethylacetate as eluent. This gave two products: 120 mg of residue which was subjected to flash chromatography using min the mixture was washed with water, dried ( ${
m Na}_2{
m SO}_4$ ) and  $(BnOOC-CH_2)_2-(R)Cgl-Pro-Pig(Z)_2$  which eluted first from the solvent evaporated in vacuo to give 729 mg of a in 6.4 ml acetonitrile and heated to reflux. After 1 h 20 mmol) H-(R)Cgl-Pro-Plg(Z) $_2$  and 0.531 g (2.996 mmol)  $_{\rm K_2CO_3}$ startingmaterials was added to a mixture of 0.64 g (0.999 0.298 g (0.999 mmol) Bnooc-CH<sub>2</sub>-OTf (see preparation of

ring, wich exhibit a broad peak ranging from 3.5 to 4.6 pronounced for the 2- and 6-CH2 groups of the piperidin piperidin ring, are selectively broader due to an  $^{
m 1}H ext{-NMR}$  (300 MHz, CDCl $_3$ ); Some signals, especially in the intramolecular exchange process. This is especially

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7.6 (m, 15H), 10.52 (bs, 1H). 1H), 3.3-3.5 (m, 4H), 4.5-4.61 (d, 1H), 5.1 (s, 6H), 7.1-2H), 2.98-3.06 (m, 1H), 3.06-3.15 (d, 1H), 3.15-3.25 (m, 6 0.94-2.27 (m, 19H), 2.28-2.43 (m, 1H), 2.8-2.98 (m,

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(iv)  $Hooc-cH_2-(R)cgl-Pro-Pig \times 2 Hcl$ 

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M hydrochloric acid followed by freeze drying gave the conversion to hydrochloric acid salt by dissolving in 1 title compound. Removal of the solvent and excess  $\mathrm{NH_4OAc}$  by freeze drying, purified on RPIC using  $m CH_3CN/0.1~M~NH_4OAc~15/85~as~eluent.$ Pro-Pig x 2 HCl. This crude material (79 % purity) was in vacuo and freeze drying gave 95 mg of HOOC-CH2-(R)Cglmillipore filter followed by evaporation of the solvent Removal of the catalyst by filtration on hyflo and acid, 10 ml ethanol (99.5 %) and 180 mg 5 % Pd/C for 2 h. hydrogenated in the presence of 0.88  $\pm$ 1 M hydrochloric 142 mg (0.176 mmol) BnOOC-CH $_2$ -(R)Cgl-Pro-Pig(Z) $_2$  was

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<sup>1</sup>H-NMR (500 MHz, D<sub>2</sub>0); & 1.1-1.35 (m, 6H), 1.63-2.14 (m, 13Н), 2.26-2.36 (ш, 1Н), 3.01-3.23 (ш, 4Н), 3.49-3.62 2H), 4.18-4.32 (d, 1H), 4.37-4.5 (m, 1H). (qvart., 2H), 3.62-3.77 (m, 2H), 3.77-3.88 (apparent d,

#### Example 5

### H-(R)Cha-Aze-Pig x 2 HCl

10 (i) Boc-(R)Cha-Aze-Pig(Z)2

20 5 of the title compound.  $\mathrm{CH_2Cl_2/MeOH}$  (97/3 followed by 95/5) to yield 43 mg (24 %) chromatography (36 g  $\mathrm{Sio}_2$ ) using a stepwise gradient of an oil. The crude product was purified by flash Filtration and evaporation of the solvent gave 141 mg of ml NaHCO3, 3 x 5 ml H2O, 1 x 5 ml brine and dried (MgSO4). organic phase was washed with 3 imes 5 ml 1 M KHSO4, 1 imes 5 and the residue was dissolved in 70 ml EtoAc and the for 20 h at room temperature. The solvent was evaporated materials) and 115 mg (0.944 mmol) DMAP in 5 ml  $\mathrm{CH_{3}CN}$  was added 50 mg (0.260 mmol) EDC and the reaction was stirred 100 mg (0.236 mmol) H-Pig(Z) $_2$  (See preparation of starting (R)Cha-Aze-OH (See preparation of starting materials), To a well stirred mixture of 86 mg (0.243 mmol) Boc-

### (ii) H-(R)Cha-Aze-Pig(Z)2

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35 30 solvent was evaporated to give 38 mg, wich was subjected in ethyl acetate as eluent to give 28 mg of the desired to flash chromatography using 10 % NH3-saturated methanol was washed with water and brine and dried  $(\mathrm{Na_2SO_4})$  . The added. The phases were separated and the organic phase in vacuo and ethyl acetate and 0.1 M NaOH-solution was ethylacetate during 5 minutes. The solvent was evaporated (0.0565 mmol) Boc-(R)Cha-Aze-Pig(Z)2 in 10 ml of Hydrogen chloride was bubbled through a mixture of 43 mg

product.

ring, wich exhibit a broad peak ranging from 3.7 to 4.5  $^{
m H-NMR}$  (300 MHz, CDCl $_3$ ); Some signals, especially in the piperidin ring, are selectively broader due to an intramolecular exchange process. This is especially pronounced for the 2- and 6-CH $_2$  groups of the piperidin

8 0.75-1.85 (m, 18H), 2.35-2.53 (m, 1H), 2.62-2.78 (m, 1H), 2.8-3.0 (m, 2H), 3.0-3.28 (m, 2H), 3.28-3.37 (m, 1H), 3.97-4.18 (m, 2H), 4.8-4.9 (m, 1H), 5.1 (s, 4H), 7.2-7.45 (m, 9H), 8.05-8.15 (m, 1H).

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(iii) H-(R)Cha-Aze-Pig x 2 HCl 15

Removal of the catalyst by filtration and evaporation in freeze drying gave 12 mg (60 %) of H-(R)Cha-Aze-Pig ethanol (99.5 %) and 0.13 ml hydrogen chloride (1 N) was hydrogenated in the presence of 35 mg 5 % Pd/C for 4 h. vacuo of the solvent followed by dissolving in water and 28 mg (0.042 mmol) H-(R)Cha-Aze-Pig(z) $_2$  dissolved in 2 ml dihydrochloride.

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ring, wich exhibit a broad peak ranging from 3.7 to 4.5 in the piperidin ring, are selectively broader due to an intramolecular exchange process. This is especially pronounced for the 2- and 6-CH $_2$  groups of the piperidin <sup>1</sup>H-NMR (500 MHz, 300 K, CD<sub>3</sub>OD); Some signals, especially

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3.0-3.12 (t, 2H), 3.12-3.23 (d, 2H), 3.85-3.95 (d, 2H), 6 0.75-2.1 (m, 18H), 2.2-2.35 (m, 1H), 2.62-2.75 (m, 1H), 3.95-4.0 (dd, 1H), 4.15-4.23 (m, 1H), 4.35-4.42 (m, 1H), 4.72-4.78 (m, 1H).

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13C-NMR (75 WHz, CD3OD): guanidine: 6 157.6; carbonyl carbons: 6 170.0 and 172.6.

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Example 52

HOOC-CH2-(R) Cgl-Aze-Pac x 2 HCl

(i) Boc-(R)Cgl-Aze-Pac(Z)

evaporated. Plash chromatography using ethyl acetate followed by ethyl acetate/methanol 98/2 as eluents gave 0.25 g (30%) of the title compound as a mixture of 1,4cis- and trans-products with respect to the Pac part of (aq) and NaHCO $_3$  (aq), dried (Na $_2$ SO $_4$ ), filtered and and 0.67 g (5.5 mmol) of DMAP in 5 ml of acetonitril was added 0.27 g of EDC at 0°C. The mixture was stirred at room temperature over night and subsequently diluted To a solution of 0.47 g (1.4 mmol) of Boc-(R)Cgl-Aze-OH (See preparation of starting materials), 0.40 g (1.4 mmol) of H-Pac(2)(See preparation of starting materials) with ethyl acetate. The solution was washed with  $ext{KHSO}_{f q}$ 

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2 H), 3.0-3.4 (m, 2 H), 3.85 (m, 1 H), 4.14 (m, 1 H), 4.33 JH-NMR (500 MHz, CDCl<sub>3</sub>) : 6 0.8-2.0 (m, 29 H; thereof 1.45 (s, 9 H)), 2.15 and 2.34 (m, 1 H, isomers), 2.45-2.7 (m, (m, 1 H), 4.85 (m, 1 H), 4.98 (m, 1 H), 5.04 (s, 2 H), 7.25-7.45 (m, 5 H), 7.8-7.9 (m, 1 H), 9.2-9.5 (m, 1 H). 52

the molecule.

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(ii) H-(R)Cgl-Aze-Pac(Z) x HCl

Boc-(R)Cgl-Aze-Pac(2), 0.25 g (0.41 mmol), was dissolved HCl (g) was bubbled through for 5 min and the solvent in 100 ml of ethyl acetate and cooled in an ice bath. was evaporated.

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1 H), 3.05 and 3.37 (multiplets, 0.6 H and 0.4 H 1H-NWR (300 MHz, MeOD) : 6 0.8-2.0 (m, 22 H), 2.05-2.35 respectively, isomers), 3.15-3.3 (m, 1 H), 4.05-4.2 (m, (m, 1 H), 2.4-2.55 (m, 1 H), 2.6-2.75 (M, 1 H), 3.00 (d,

2 H), 4.88 (dd, 1 H), 5.11 (s, 2 H), 7.2-7.45 (m, 5 H), 8.0-8.15 (m, 1 H).

## (iii) BnO<sub>2</sub>C-CH<sub>2</sub>-(R)Cgl-Aze-Pac(Z)

A mixture of 0.17 g (0.33 mmol) of H-(R)Cgl-Aze-Pac(Z) x HCl, 0.11 g (0.37 mmol) of benzyl triflyloxyacetate and 0.14 g (1.0 mmol) of K<sub>2</sub>CO<sub>3</sub> in 5 ml of acetonitrile was stirred at room temperature for 3 days. The crude material was flash chromatographed with EtOAc/CH<sub>2</sub>Cl<sub>2</sub>/MeOH 95/20/5. Yield: 70 mg (32 %).

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1H-NMR (500 MHz, CDCl<sub>3</sub>): & 0.85-2.3 (m, 20 H), 2.48 (m, 1 H), 2.63 (m, 1 H), 2.87 (m, 1 H), 3.05-3.25 (m, 1 H), 3.25-3.35 (m, 2 H), 3.38 (dd, 1 H), 3.95 (m, 1 H), 4.08 (m, 1 H), 4.88 (m, 1 H), 5.1-5.2 (m, 4 H), 5.9-6.3 (m, 1 H), 7.25-7.5 (m, 10 H), 8.00 and 8.08 (broad triplets, 1 H, isomers).

## (iv) $Ho_2C-CH_2-(R)Cgl-Aze-Pac \times 2 HCl$

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BnO<sub>2</sub>C-CH<sub>2</sub>-(R)Cgl-Aze-Pac(Z), 70 mg (0.11 mmol), was dissolved in 5 ml of ethanol, and 5% Pd/C and 0.1 ml of conc. HCl were added. The mixture was hydrogenated at etmospheric pressure for 1 h. After filtration and evaporation the product was purified through preparative RPLC using 0.1 M NH<sub>4</sub>OAC/CH<sub>3</sub>CN 4/1 as eluent. After change of salt to the hydrochloride and freeze drying 1,4-cis- and trans-isomers with respect to the Pac part of the molecule. Yield: 40 mg (74%).

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<sup>1</sup>H-NMR (500 MHz, D<sub>2</sub>0) & 1.1-2.1 (m, 20 H), 2.32 (m, 1 H), 2.52 (m, 1 H), 2.63 (m, 1 H), 2.72 (m, 1 H), 3.1-3.3 (m, 1 H), 3.40 (m, 1 H), 3.8-3.95 (m, 2 H), 4.04 (d, 1 H), 4.39 (m, 1 H), 4.93 (m, 1 H).

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 $^{13}\text{C-NMR}$  (125 MHz,  $D_2\text{O})$  amidine and carbonyl carbons: § 167.7, 172.0, 174.9 and 175.2.

#### Example 53

### H-(R)Cha-Pro-Pac x 2 HCl

### (i) Boc-(R)Cha-Pro-Pac(Z)

20 15 10 which was purified by flash chromatography using ethyl water, and sodium hydrogen carbonate solution and dried acetate as eluent to give 196 mg (27%) of the title  $({
m MgSO}_{f q})$  . Removal of the solvent in vacuo gave a residue water. The organic phase was washed with acetic acid, and the residue was diluted with ethyl acetate and temperature for 2 h. The solvent was removed in vacuo reaction mixture was stirred at 0°C for 1 h and at room materials), and 0.55 g DMAP in 7 ml acetonitrile. The Boc-(R)Cha-Pro-OH(See preparation preparation of starting materials), 0.4 g (1.1 mmol) solution of 0.4 g (1.1 mmol) H-Pac(Z) x 2 HCl (See 211 mg (1.1 mmol) EDC was added at 0°C to a stirred 얁 starting

### 25 (ii) H-(R)Cha-Pro-Pac(Z)

Hydrogen chloride was bubbled through a solution of 196 mg Boc-(R)Cha-Pro-Pac(Z) in 25 ml ethyl acetate. After 10 minutes the reaction mixture was diluted with methylene chloride and sodium hydroxide solution was added. The aqueous phase was extracted several times with methylene chloride and the combined organic phases were dried (K<sub>2</sub>CO<sub>3</sub>) and the solvent was removed in vacuo to give 86 mg (52%) of the title compound.

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The title compound was prepared by hydrogenation of H-(R)Cha-Pro-Pac(Z) in ethanol in the presence of 10% Pd/C.

1,4-trans isomers in the Pac part of the molecule);  $\delta$ 1.15-1.3 (q), 1.6-1.85 (m), 1.9-2.0 (m), 2.0-2.1 (d), 2.1-2.15 (m), 2.15-2.2 (m), 2.65-2.7 (m), 2.7-2.8 (m), H-NMR (300 MHz, D20;A ca: 1:1 mixture of 1,4-cis- and 2.95-3.0 (d), 3.15-3.2 (d), 5.4 (s), 7.45-7.55 (m).

Example 54

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H-(R)cgl-Ile-Pab x 2 HCl

(i) Boc-(R)Cgl-Ile-Pab(Z) 12

phase was washed with 2  $\times$  30 ml NaHCO $_3$ (saturated), 2  $\times$ Evaporation followed by flash chromatography using extracted with 2  $\times$  50 ml EtOAc and the combined organic 50 ml 0.2 M HCl, 1 x 50 ml Brine and dried(MgSO4).  $CH_2Cl_2/THF$  (85/15) as eluent gave 510 mg (24 %) of the temperature and left for 60 h. The  $\mathrm{CH_{3}CN}$  was removed by evaporation and the residue was poured out in 100 m 
m Iwater (a yellow precipitate was formed). The mixture was Ile-OH (See Preparation of starting materials), 1.12 g  $\mathrm{CH_3CN/DMF}$  (1/1) was added 0.75 g (3.9 mmol) EDC at + 5°C. The reaction mixture was allowed to reach room (3.9 mmol) H-Pab(Z) (See Preparation of starting materials) and 1.76 g (14.4 mmol) DMAP in 50 ml To a stirred mixture of 1.33 g (3.6 mmol) Boc-(R)Cgltitle compound. ဓ္ဗ

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(11) H-(R)Cgl-Ile-Pab(Z)

530 mg Boc-(R)Cgl-Ile-Pab(Z) was dissolved in 14 ml  $CH_2Cl_2/TFA$  (2.5/1) and stirred for 2 h at room temperature. Evaporation of the solvent followed by 35

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flash chromatography using  ${
m CH_2Cl_2/MeOH\,(NH_3-saturated)}$ (95/5) as eluent gave the title compound.

(iii) H-(R)Cgl-Ile-Pab x 2 HCl

pressure for 6 h. Addition of 2 g activated charcoal and evaporation of the solvent and freeze drying from water gave 50 mg (89%) of the title compound as a white 75 mg (0.14 mmol) H-(R)Cgl-Ile-Pab(Z) was hydrogenated over 10 % Pd/c in 5 ml EtOH, which contained an excess HCl(g) to give the dihydrochloride, at atmospheric 20 ml EtOH followed by filtration through celite, powder. 2

1H-NWR(500 MHz, MeOD): 6 0.90 (t, 3H), 0.94 (d, 3H), 1.1-2.0 (m, 14H), 3.83 (bs, 1H), 4.26 (d, 1H), 4.50 (m, 2H), 7.57 (bd, 2H), 7.78 (bd, 2H).

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Example 55

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H-(R)Cgl-Aze-Pab

(see Example 1 (ii)) over 5 % Pd/C in 6 ml  $BtOH/H_2O$  at atmospheric pressure for 6 h followed by filtration of the catalyst, evaporation of the solvent and freeze drying from water gave 200 mg (89 %) of the title Hydrogenation of 257 mg (5.08 mmol)  $H^-(R)$ Cgl-Aze-Pab(Z) compound.

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2.70 (m, 1H), 3.30 (m, 1H), 3.75 (m, 1H), 4.30 (m, 1H), 4.45 (m, 1H), 4.55 (m, 2H), 7.60 (m, 2H), 7.77 (m, 2H). 1H-NMR (500 MHz, D<sub>2</sub>0): 6 1.0-2.0 (m, 11H), 2.25 (m, 1H), ဓ္က

MS m/z 372 (M + 1).

#### Example 56

## HOOC-(R, S)CH(Me)-(R)Cha-Pro-Pab x HOAc

## (i) BnOOC-(R,S)CH(Me)-(R)Cha-Pro-Pab(Z)

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followed by flash chromtography using  $ext{CH}_2 ext{Cl}_2/ ext{MeOH}$  9/1 as with water and dried (MgSO $_4$ ). Evaporation of the solvent eluent gave 150 mg (46%) of the title compound. 20 h. The mixture was diluted with  $\mathrm{CH_2Cl_2}$  , extracted added and the mixture was stirred at roomtemperature for slowly. 200 mg (1.45 mmol) of potassium carbonate was Startingmaterials) dissolved in 3 ml  $\text{CH}_2\text{Cl}_2$  was added dissolved in 5 ml  $ext{CH}_2 ext{Cl}_2$  was cooled to -10°C and 150 mg (0.48 mmol) of TfOCH2COOBn (See Preparation of 0.250 g (0.47 mmol) H-(R)Cha-Pro-Pab(Z) (See Example 15)

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## (ii) HOOC-(R,S)CH(Me)-(R)Cha-Pro-Pab x HOAc

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eluent, gave 35 mg (37%) of the title compound. purification by RPLC, using  $CH_3CN/0.1\ M\ NH_4OAc\ 1/4\ as$ catalyst, evaporation of the solvent followed by athmospheric pressure for 4 h. Filtration of the was hydrogenated over 50 mg 5 % Pd/C in 20 ml EtOH at 150 mg (0.2 mmol) BnOOC-(R,S)CH(Me)-(R)Cha-Pro-Pab(Z)

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the HOD line, 6H), 7.55 (d, 2H), 7.75 (d, 2H). (m, 1H), 4.35-4.6 (m, 3H), 4.9 (m, partially hidden by 5H), 1.5 (m, 1H), 1.6-1.8 (m, 6H), 1.9-2.1 (m, 6H), 2.25  $^{1}\text{H-NMR}$  (500 MHz, MeOD): & 1.00 (m, 1H), 1.20-1.45 (m, (m, 1H), 3.25 (m, 1H), 3.5 (m, 1H), 3.85 (m, 1H), 4.15

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#### Example 57

## MeOOC-CH<sub>2</sub>-(R)Cgl-Aze-Pab x 2 HCl

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(i) MeOOC-CH<sub>2</sub>-(R)Cgl-Aze-Pab(Z)

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G and the last time diethylether/MeOH(NH3-saturated) CH<sub>2</sub>Cl<sub>2</sub>/THF/MeOH (16/4/1), then CH<sub>2</sub>Cl<sub>2</sub>/THF(2%NH<sub>3</sub>) (8/2) (95/5) as eluent. This gave 0.324 g (67 %) of the title chromatography residue that was three times subjected to flash solvent was evaporated in vacuo to give 0.51 g of a washed with water and brine, dried, filtered and the over night. More  $ext{CH}_2 ext{Cl}_2$  was added and the mixture was  $ext{CH}_2 ext{Cl}_2$  (totally 4.3 ml) at roomtemperatur, and stirred Pab(Z) (See Example 1), 0.894 g (5.04 mmol)  $K_2CO_3$  in added to a mixture of 0.425g (0.841 mmol) H-(R) Cgl-Azestarting materials) was dissolved in  $ext{CH}_2 ext{Cl}_2$  and slowly 0.186 g (0.841 mmol) TfO-CH $_2$ -COOMe (See preparation of on silica gel using, first

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## (ii) MeOOC-CH<sub>2</sub>-(R)Cgl-Aze-Pab x 2 HCl

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20 twice gave 178 mg (91 %) of the title compound. by evaporation of the solvent in vacuo and freeze drying by filtration on cellite and millipore filter followed MeOH and 300 mg Pd/C for 2 h. Removal of the catalyst hydrogenated in the presence of 1.14 ml 1 N HCl, 6.5 ml 220 mg (0.38 mmol) MeOOC-CH<sub>2</sub>-(R)Cgl-Aze-Pab(Z) was

4.5 (d, 1H), 4.36-4.42 (t, 2H), 4.59 (s, 2H), 4.99-5.04 1H), 2.68-2.8 (m, 1H), 3.86 (s, 3H), 4.1 (s, 2H), 4.1-2H), 1.81-1.9 (m, 3H), 1.97-2.1 (m, 1H), 2.29-2.4 (m, <sup>1</sup>H-NMR (500 MHz, D<sub>2</sub>0); & 1.12-1.4 (m, 5H), 1.68-1.81 (m, (m, 1H), 7.65-7.7 (d, 2H), 7.8-7.85 (d, 2H).

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δ 146.78, 167.68, 168.15, 172.29. 13C-NMR (75 MHz, MeOD): amidine and carbonyl carbons; 30

#### Example 58

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Etooc-cH2-(R)cgl-Aze-Pab x 2 HCl

(i) EtOOC-CH2-(R)Cgl-Aze-Pab(Z)

 $CH_2Cl_2$  (totally 4 ml) cooled on an ice-bath. After 2 h the ice-bath was removed and stirring was continued at roomtemperature for 2 hours. More  $extsf{CH}_2 extsf{Cl}_2$  was added and filtered and the solvent was evaporated in vacuo to give 0.51 g of a residue that was subjected to flash chromatography using diethylether/MeOH( $\mathrm{NH_3} ext{-saturated}$ ) (95/5) as eluent. This gave 0.387  $\mathfrak g$  (75 %) of the title Pab(Z) (See Example 1) and 0.931 g (5.26 mmol)  $\rm K_2CO_3$  in the mixture was washed with water and brine, dried, 0.208 g (0.876 mmol) TfO-CH<sub>2</sub>-COOEt (See preparation of starting materials) was dissolved in  ${
m CH}_2{
m Cl}_2$  and slowly added to a mixture of 0.443 g (0.876 mmol) H-(R)Cgl-Aze-

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(ii) Etooc-CH<sub>2</sub>-(R)Cgl-Aze-Pab  $\times$  2 HCl

compound.

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2 eqvivalents of 1 N HCl was added, and freeze drying 390 mg Pd/C for 5 h. Removal of the catalyst by filtration on cellite and millipore filter, followed by evaporation of the solvent in vacuo and freeze drying twice, gave 281 mg (88 %) of EtooC-CH<sub>2</sub>-(R)Cgl-Aze-Pab. 395 mg (0.668 mmol) Etooc-CH<sub>2</sub>-(R)Cgl-Aze-Pab(Z) was hydrogenated in the presence of 12 ml EtOH (99.5 %) and three times gave 288 mg (81 %) of the title compound.

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6H), 2.15-2.33 (m, 1H), 2.58-2.79 (m, 1H), 3.89-4.0 (m, 3H), 4.2-4.33 (m, 3H), 4.33-4.44 (m, 1H), 4.44-4.66 (m, 2H), 4.91 (m, 1H (partially hidden by the H-O-D  $^{1}_{H-NMR}$  (500 MHz,  $^{0}_{2}$ 0); 6 1.05-1.48 (m, 8H), 1.6-2.05 (m, signal)), 7.54-7.63 (d, 2H), 7.72-7.84 (d, 2H).

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Example 59

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"Bucoc-CE2-(R) cgl-Ase-Pab x HOAC

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(i) "Buooc-CH2-(R)Cgl-Aze-Pab(Z)

COO<sup>n</sup>Bu as alkylating agent. The crude product was purified by flash chromatography twice, first using  ${
m CH_2Cl_2/MeOH}$  (95/1) as eluent and then  ${
m CH_2Cl_2/i-}$ propylalcohol (90/7) to give 324 mg (47 %) of the title Prepared in the same way as described for  $^{\mathrm{n}}\mathrm{HexOOC}\text{-CH}_{2}\text{-}$ (R)Cgl-Aze-Pab(Z) (See Example 60 (1)) using  $TCO-CH_2$ combound.

(ii) "Buooc-CH2-(R)Cgl-Aze-Pab x HOAc

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The deprotection was done according to the procedure described in Example 57 (ii). The crude material was purified on RPLC using  $\mathrm{CH_3CN}$  (30 %) in 0.05 M  $\mathrm{NH_4OAc}$  and 0.05 M HOAc as eluent to give 100 mg (53 %) of the title combound.

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(m, 1H), 2.58-2.8 (m, 1H), 3.7-5.0(m, 10H), 4.88-5.0 <sup>1</sup>H-NYR (300 MHz, MeOD); 6 0.85-2.1 (m, 18H), 2.15-2.37 (partially hidden by the H-O-D signal)), 7.46-7.65 (d, H), 7.71-7.88 (d, 2H). 20

13C-NMR (75 MHz, MeOD): amidine and carbonyl carbons; 6 146.8, 168.12, 168.2, 172.2. 25

Example 60

"HexOOC-CH2-(R) CG1-Are-Pab x 2 HC1 30

(i) "HexOOC-CH2-(R) Cgl-Aze-Pab(Z)

Pab(2) (See Example 1), 1.463 g (8.25 mmol)  $K_2 CO_3$  in starting materials) was dissolved in  ${\rm CH}_2{\rm Cl}_2$  and slowly 0.402 g (1.375 mmol) TfO-CH $_2$ -COO $^{\mathrm{h}}$ Hex (See Preparation of added to a mixture of 0.695 g (1.375 mmol) H-(R)Cgl-Aze-35

saturated) (95/5), and then  $\mathrm{CH_2Cl_2/MeOH(NH_3-saturated)}$ chromatography, first using diethylether/MeOH(NH $_3-$ 0.828 g of a residue, wich was twice subjected to flash compound. (95/5) as eluent. This gave 0.42 g (47 %) of the title filtered and the solvent was evaporated in vacuo to give temperature for 45 minutes. More  $\mathrm{CH_2Cl_2}$  was added and the mixture was washed with water and brine, dried, bath was removed and stirring was continued at room  $\mathrm{CH_2Cl_2}$  (totally 4 ml) at <-10°C. After 1 h the  $\mathrm{CO_2}\text{-ice-}$ 

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## (ii) ThexOOC-CH<sub>2</sub>-(R)Cgl-Aze-Pab x 2 HCl

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and freeze drying twice, gave 287 mg (79 %) of the title filter, followed by evaporation of the solvent in vacuo 1.7 ml 1 N HCl, 12 ml MeOH and 340 mg Pd/C. Removal of hydrogenation was completed in 4 h in the presence of the catalyst by filtration on cellite and millipore for 1.5 h did not give complete de-protection. The Aze-Pab(Z) in the presence of 12 ml THF and 400 mg Pd/C Hydrogenation of 400 mg (0.617 mmol)  $^{n}$ HexOOC-CH $_{2}$ -(R)Cgl-

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hidden by the H-O-D signal)), 7.52-7.69 (d, 2H), 7.75-7.9 (d, 2H). (m, 3H), 4.37-4.7 (m, 3H), 4.88-5.0 (m, 1H (partialyy (m, 1H), 2.61-2.81 (m, 1H), 3.93-4.15 (m, 3H), 4.15-4.37 1H-NMR (300 MHz, MeOD); & 0.8-2.13 (m, 22H), 2:13-2.31

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13C-NMR (75 MHz, MeOD): amidine and carbonyl carbons; 6 146.84, 167.67, 167.84, 172.17.

Example 61

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H-(R)Cgl-Pro-Pac x 2 HCl

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(i) Boc-(R)Cgl-Pro-Pac(Z)

giving MS m/z = 626 (M + 1) were isolated. subsequently by RPIC. Two fractions (51 mg and 150 mg) using 10 \$ methanol in methylene chloride as eluent, and residue was first purified by flash chromatography, over night. The solvent was removed in vacuo and the reaction mixture was allowed to reach room temperature 1.078 g (8.8 mmol) DMAP in 12.5 ml acetonitrile. The Preparation of startingmaterial), 714 mg (2.0 mmol) H-Pac(Z) imes 2 HCl (See Preparation of startingmaterial) and solution of 708 mg (1.95 mmol) of Boc-(R)Cgl-Pro-OH (See 377 mg (1.97 mmol) EDC was added at 0°C to a stirred

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15 (ii) H-(R)Cgl-Pro-Pac(Z)

the product. was added and the organic phase was separated and dried acetate. After 15 minutes 10 % sodium carbonate solution  $(\mathrm{K}_2\mathrm{CO}_3)$  . Evaporation of the solvent gave 71 mg (61 %) of Hydrogen chloride was bubbled into a solution of 141 mg (0.22 mmol) Boc-(R)Cgl-Pro-Pac(Z) in 50 ml ethyl

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(iii) H-(R)Cgl-Pro-Pac x 2 HC]

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and the solvent was removed in vacuo. The residue was Freeze drying yielded 38 mg (58 %) of the title compound dissolved in 50 ml water and 0.6 g 1M hydrochloric acid. pressure for 2 h. The catalyst was removed by filtration hydrogenated at room temperature and atmospheric a small spatula of 10 % Pd/C in 10 ml of ethanol was A mixture of 71 mg (0.14 mmol) H-(R)Cgl-Pro-Pac(Z) and

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MS m/z 392 (M + 1)

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Example 62

HOOC-CH2-(R) Cha-Pro-Pac x HOAC

(i) BnOOC-CH<sub>2</sub>-(R)Cha-Pro-Pac(Z)

using ethyl acetate/methylene chloride/methanol 95:20:5 filtered and the solvent was removed in vacuo to give a residue which was subjected to flash chromatography materials) in 3 ml of methylene chloride was stirred at room temperature over night. The reaction mixture was A mixture of 84 mg (0.15 mmol) H-(R)Cha-Pro-Pac(Z) (See and 47 mg of TfOCH $_2$ -COOBn (See Preparation of starting Example 53 (11)), one spatula of potassium carbonate, as eluent. 29 mg of the desired product was isolated.

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(11) HOOC-CH2-(R)Cha-Pro-Pac x HOAc

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catalyst followed by removal of the solvent and A mixture of 29 mg BnOoC-CH $_2$ -(R) Cha-Pro-Pac(Z) and 37 mg temperature and atmospheric pressure. Filtration of the of 10 % Pd-C in 5 ml etanol was stirred for 4 h at room purification by RPLC gave the desired compound.

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MS m/z = 464 (M + 1). 22

Example 63

HOOC-CH2-CH2-(R) CG1-Pro-Pac

(i) Bnooc-CH2-CH2-(R) Cgl-Pro-Pac(Z)

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acrylate, and 280  $\mu l$  (2 mmol) triethyl amine in 1 ml ethanol was kept at room temperature for 3 days. Removal of the solvent followed by purification by HPLC gave 18 (See Example 61 (11) ), 124 mg (0.76 mmol) benzyl A solution of 0.35 g (0.64 mmol) H-(R)Cgl-Pro-Pac(Z)

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mg (4 %) of the title compound.

(ii) HOOC-CH2-CH2-(R)Cgl-Pro-Pac

a small spatula of 10 % Pd/C was hydrogenated for 2 h Filtration followed by removal of the solvent in vacuo and dissolution in water and freeze drying gave 7 mg (78 A mixture of 18 mg BnoOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cgl-Pro-Pac(Z) and at room temperature and atmospheric pressure in EtOH. %) of the title compound. MS m/z = 464 (M + 1).

Example 64

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(i) Boc-(R)Cha-Aze-Pac(Z)

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was partitioned between ethyl acetate and water. The agueous phase was extracted once more with ethyl acetate and brine and then dried (sodium sulphate). Evaporation acetonitrile was mixed at  $0^{\circ} C$  with a solution of 0.26g (1.4 mmol) EDC in 15 ml acetonitrile. The reaction mixture was kept at room temperature over night and the solvent was subsequently removed in vacuo. The residue and the combined organic phases were washed with sodium hydrogen sulphate solution, sodium carbonate solution, startingmaterial), and 0.67 g (5.5 mmol) DWAP in 20 ml of the solvent gave 0.54 g (63 %) of the title compound. 0.5 g (1.41 mmol) Boc-(R)Cha-Aze-OH(See Preparation of Preparation of startingmaterial of  $H ext{-Pac}(2)$  x 2 HCl), A solution of 0.4 g (1.38 mmol) H-Pac( $\mathbb{Z}$ )

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(ii) H-(R)Cha-Aze-Pac(Z)

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solution was kept in the refrigerator over night and Hydrogen chloride was bubbled into a solution of 0.54 g (0.9 mmol) Boc-(R)Cha-Aze-Pac(Z) in ethyl acetate. The

solvent gave 0.35 g (77 %) of the product. and brine and dried (sodium sulphate). Removal of the with aqueous sodium hydrogen carbonate solution, water was dissolved in ethyl acetate. The solution was washed the solvent was then removed in vacuo and the residue

## (iii) BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cha-Aze-Pac(2)

15 10 chloride as eluent to yield 150 mg ( 66 %) of the title flash chromatography, using 10 % methanol in methylene solvent in vacuo gave a residue which was purified by brine. Drying (sodium sulphate) and removal of the The solution was washed with potassium hydrogen sulphate at room temperature for 60 h. The solvent was removed solution and sodium hydrogen carbonate solution and in vacuo and the residue was dissolved in ethyl acetate. 53 mg (0.33 mmol) benzyl acrylate in ethanol was kept A solution of 180 mg (0.33 mmol) H-(R)Cha-Aze-Pac(Z) and

## (iv) HOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cha-Aze-Pac x 2 HCl

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dried to give 30 mg (33 %) of the title compound. and dissolution of the residue in water and 1.5 ml of 1.5 h at room temperature and atmospheric pressure. 1M hydrochloric acid gave a solution which was freeze Filtration followed by removal of the solvent in vacuo 67 mg of 10 % Pd-C in 10 ml ethanol was hydrogenated for A mixture of 115 mg BnOOC-CH $_2$ -CH $_2$ -(R)Cha-Aze-Pac(Z) and

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MS m/z 464 (M + 1).

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#### Example 65

HOOC-CH2-(R)Cha-Aze-Pig x 2 HC1

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(i) Boc-(R)Cha-Aze-Pig(Z)

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10 the title compound. EtOAc/MeOH 9/1 as eluent. This gave 407 mg (53 %) of evaporation of the solvents gave 0.612 g of a residue washed with saturated  ${
m Na_2CO_3}$  and brine. Repeting the which was subjected to flash chromatography using extractive procedure, drying  $(\mathrm{Na_2SO_4})$ , filtration and The phases were separated and the organic phase was evaporated in vacuo and EtOAc and 2 M  $ext{KHSO}_4$  was added roomtemperature over night. The solvent was starting materials) and 0.604 g (4.94 mmol) DMAP in 13.5 ml DMF. The temperature was allowed to rise to (1.236 mmol) H-Pig(Z)  $\times$  HCl (See Preparation of mixture of 0.473 g (1.236 mmol) Boc-(R)Cha-Aze-OH 0.249 g (1.298 mmol) EDC was added at <-15°C to a (See Preparation of starting materials), 0.404 g

#### (ii) H-(R)Cha-Aze-Pig(Z)

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25 20 %) of the title compound. water and brine, dried  $(Na_2SO_4)$ , filtered and the solvent was evaporated in vacuo to give 336 mg (100 were seperated and the organic layer was washed with and EtOAc and saturated  $\mathrm{Na_2CO_3}$  was added. The phases roomtemperature. The solvent was removed in vacuo minutes on an ice-bath, and for 30 minutes at dissolved in 24.4 ml of TFA/CH<sub>2</sub>Cl<sub>2</sub> 1/4, stirred for 30 0.4 g (0.638 mmol) of Boc-(R)Cha-Aze-Pig(Z) was

## (iii) BnOOC-CH<sub>2</sub>-(R)Cha-Aze-Pig(Z)

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was evaporated, EtOAc was added, and the mixture was washed with water, dried ( $Na_2SO_4$ ), filtered and the solvent was evaporated in vacuo to give 346 mg of a 60°C on an oilbath. After 1 h 45 minutes the solvent mixture of 0.296 g (0.562 mmol) H-(R)Cha-Aze-Pig(Z) and 0.171 g (1.236 mmol)  $K_2CO_3$  in 6 ml  $CH_3CN$  heated to 89 ml (0.562 mmol) BnOOC-CH $_2$ -Br was slowly added to a

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using CH<sub>2</sub>Cl<sub>2</sub>/THF/MeOH (8/2/1) as eluent. This gave 297 residue which was subjected to flash chromatography mg (78 %) of the title compound.

(1v)  $HOOC-CH_2-(R)Cha-Aze-Pig \times 2 HCl$ 

hydrogenated in the presence of 1.7 ml 1 N HCl, 10 ml  $\,$ EtcH (99.5 %) and 300 mg Pd/C for 2 h. Removal of the vacuo and freeze drying twice gave 166 mg (88 %) of 243 mg (0.36 mmol) Bnooc-CH<sub>2</sub>-(R)Cha-Aze-Pig(Z) was filter followed by evaporation of the solvent in catalyst by filtration on cellite and millipore the title compound

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(m, 1H), 2.52-2.76 (m, 1H), 2.82-3.2 (m, 4H), 3.46-JH-NMR (500 MHz, D<sub>2</sub>0); 6 0.6-1.9 (m, 18H), 2.1-2.27 3.61 (m, 1H), 3.61-3.81 (m, 2H), 3.81-4.0 (m, 2H), 4.0-4.24 (m, 2H), 4.24-4.4 (m, 1H).

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Example 66 20

HOOC-CH2-(R) Cha-Pro-Pig x 2 HCl

(i) Boc-(R)Cha-Pro-Pig(Z)

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layer was dried ( $Na_2SO_4$ ), filtered and evaporated. The using a stepwise gradient of  ${
m CH_2Cl_2/MeOH~(100/0,~97/3)}$ (see Preparation of starting materials) in 5 ml  ${\rm CH_2Cl_2}$ (3.8 mmole) DMAP, 0.310 g (0.95 mmole) H-Pig(Z) x HCl in ethyl acetate. The organic phase was washed twice To a mixture of 0.3495 g (0.95 mmole) Boc-(R)Cha-Promixture was evaporated and the residue was dissolved OH (See Preparation of starting materials), 0.464 g crude product was purified by flash chromatography was added 0.192 g (1 mmole) of EDC and the mixture with 0.3 M KHSO4 and once with brine. The organic was stirred over night at room temperature. The 35 30

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95/5, 90/10) as eluent to yield 307 mg of the title combound.

(ii) H-(R)Cha-Pro-Pig(Z)

solvent was evaporated and the residue was dissolved in CH<sub>2</sub>Cl<sub>2</sub>. The organic layer was washed twice with 0.2 with  ${\rm CH_2Cl_2}$  and the combined organic layer was dried dissolved in 30 ml HCl saturated ethyl acetate. The M NaOH. The combined water layer was extracted once mixture was allowed to stand for half an hour. The  $(\mathrm{Na_2SO_4})$ , filtered and evaporated to yield 257 mg 0.306 g (0.48 mmole) Boc-(R)Cha-Pro-Pig(2) was (99%) of the title compound.

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(iii) Bnooc-CH2-(R)Cha-Pro-Pig(Z)

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in  $\mathrm{CH_2Cl_2}$ , washed once with water and once with brine, 0.2 g product (90% pure according to RPLC). The final mmole) of bensylbromoacetate in 6 ml acetonitrile was solvent was evaporated and the residue was dissolved  $\mathrm{CH}_2\mathrm{Cl}_2/\mathrm{MeOH}$  ( 97/3, 95/5, 90/10 ) as eluent to yield dried ( $\mathrm{Na_2SO_4}$ ), filtered and the solvent evaporated. CH2Cl2/MeOH 95/5 yielding 0.158 g (48%) of the pure Pig(2), 0.144 g (1.04 mmole)  $\rm K_2CO_3$  and 82  $\mu l$  (0.521 purification was made on a chromatotron (Harrison research, model 7924T ) on a 2mm silica plate in A mixture of 0.256 g (0.473 mmole) H-(R)Cha-Proheated to 60°C for two hours under stirring. The chromatography using a stepwise gradient of The crude product was purified by flash 30 25 20

(iv)  $HOOC-CH_2-(R)Cha-Pro-Pig \times 2 HC1$ 

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0.158 g (0.227 mmole) of BnOOC-CH2-(R)Cha-Pro-Pig(Z) was mixed with 0.075 g Pd/C (5%), 1.0 ml lN HCl-

mg (97%) of the product followed by freeze drying twice from water gave 119 through cellite and evaporation of the solvent hydrogenated at atmospheric for one hour. Filtration solution and 10 ml ethanol. The mixture was

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2H), 4.44-4.58 (m, 2H) 4H), 3.68 (m, 1H), 3.77-4.02 (m, 5H; thereof 3.98 (s, 1H), 1.60-2.20 (m, 13H), 2.39 (m, 1H), 3.07-3.32 (m, 1H-NMR (D2O, 300 MHz): 6 0.95-1.44 (m, 7H), 1.52 (m,

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carbons: & 156.5, 168.3, 169.6, 174.5  $^{13}\mathrm{C-NMR}$  (D20, 75 MHz): carbonyl- and guanidine

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HOOC-CH2-CH2-(R)Cha-Pro-Pig x 2 HC1

(i) BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cha-Pro-Pig(Z)

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eluent to yield 0.338 g (87%) of the title compound. a stepwise gradient of  $ext{CH}_2 ext{Cl}_2/ ext{MeOH}$  (95/5, 90/10 ) as research, model 7924T ) using a 2mm silica plate with product chromatographed on a chromatotron (Harrison temperature. The solvent was evaporated and the crude mixture was stirred for four days at room 66 (ii)) was dissolved in 2 ml ethanol and 90  $\mu l$ 0.297 g (0.55 mmole) H-(R)Cha-Pro-Pig(Z) (See Example (0.59 mmole) bensylacrylate was added. The reaction

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(ii) HOOC-CH2-CH2-(R)Cha-Pro-Pig x 2 HC1

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HCl-solution and 15 ml ethanol. The mixture was hydrogenated at atmospheric pressure for one hour. Pig(Z) was mixed with 0.120 g Pd/C (5%), 1.2 ml 1N Filtration of the catalyst through cellite, 0.238 g (0.227 mmole) of BnOOC-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cha-Pro-

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compound. twice from water gave 178 mg (95 %) of the title evaporation of the solvent followed by freeze drying

- 6H), 3.57 (bg, 1H), 3.67-3.87 (m, 3H), 4.25-4.43 (m, (m, 13H), 2.29 (m, 1H), 2.83 (t, 2H), 2.9-3.4 (m, <sup>1</sup>H-NMR ( $D_2$ 0, 300 MHz):  $\delta$  0.82-1.45 ( $\pi$ , 8H), 1.45-2.15
- 10  $^{13}\mathrm{C-NMR}$  (D $_2$ O, 75 MHz): carbonyl- and guanidine carbons: 6 156.3, 168.2, 174.3, 174.6

MS m/z 479 (M+1)

Example 68

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(HOOC-CH<sub>2</sub>)<sub>2</sub>-(R)Cgl-Pro-Pig x 2 HC]

30 25 20 3.8-3.9 (d, 2H), 4.07-4.22 (m, 2H), 4.22-4.35 (m, 3.28 (m, 4H), 3.58-3.70 (m, 1H), 3.7-3.8 (m, 1H), 1H), 4.38-4.5 (m, 1H). (d, 1H), 1.64-2.14 (m, 11H), 2.27-2.39 (m, 1H), 3.03- $^{1}\text{H-NMR}$  (500 MHz, D<sub>2</sub>O); 6 1.05-1.38 (m, 7H), 1.53-1.64 freeze drying gave 66 mg (90 %) of the title compound evaporation of the solvent in vacuo followed by filtration on cellite and millipore filter and 150 mg Pd/C for 4 h. Removal of the catalyst by presence of 0.75 ml 1 N HCl, 7 ml EtOH (99.5 %) and (See Example 50 (iii) ) was hydrogenated in the 120 mg (0.126) mmol (BnOOC-CH<sub>2</sub>)<sub>2</sub>-(R)Cgl-Pro-Pig(Z)<sub>2</sub>

δ 156.28, 166.73, 170.14, 174.01.  $^{13}\mathrm{C-NMR}$  (75 MHz,  $\mathrm{D_2O}$ ): amidine and carbonyl carbons;

Example 69

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HOOC-CH2-CH2-(HOOC-CH2)-(R)Cha-Pro-Pig x 2 HCl

(i) Bnooc-CH<sub>2</sub>-CH<sub>2</sub>-(Bnooc-CH<sub>2</sub>)-(R)Cha-Pro-Pig(Z)

room temperature. Evaporation of the solvent followed solution of 64 mg (0.21 mmol) TfO- $\mathrm{CH}_2\mathrm{-C00Bn}$  dissolved reflux for 30 minutes and finally left over night at in 1 ml  ${
m CH_2Cl_2}$ . The reaction mixture was left at 0°C by flash chromatography using  $\mathrm{CH_2Cl_2/MeOH}$  (97/3) as eluent afforded 65 mg (54 %) of the title compound. To a cold (ice-bath temperature) mixture of 100 mg temperature for 2 h after which it was heated to carbonate in 4 ml of  $\mathrm{CH_2Cl_2}$  was carfully added a (0.14 mmol) Bnooc-CH2-CH2-(R)Cha-Pro-Pig(Z) (See Example 67 (1)) and 80 mg (0.57 mmol) potassium for 30 minutes and then allowed to reach room

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(11) HOOC-CH2-CH2-(HOOC-CH2)-(R)Cha-Pro-Pig x 2 HCl

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evaporation of the solvent followed by freeze drying from water gave 40 mg (97 %) of the title compound as 65 mg (0.08 mmol) of Bnooc- ${
m CH_2-CH_2-(Bnooc-CH_2)-(R)\,Cha-}$ athmospheric pressure. Filtration of the catalyst (9/1) and hydrogenated over 10 % Pd/C for 3 h at Pro-Pig(Z) was dissolved in 10 ml of EtOH/1M HCl a white powder. 25

13C-NMR (125 MHz, MeOD): amidine and carbonyl carbons: & 157.5, 167.2, 169.1, 173.7 and 174.1.

Example 70

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HOOC-CH2-(R) CG1-Aze-(R, B) Itp x 2 HCl

(1) Boc-(R) Cgl-Aze-(R, S) Itp(Ts) 35 Boc-(R)Cg1-Aze-OH (See Preparation of starting

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mmol) was added and the resulting mixture was stirred separated organic layer was washed with  $\rm K_2CO_3(sat)$ , 2 at room temperature over night. The CH3CN was removed M KHSO4, brine and dried(Na<sub>2</sub>SO<sub>4</sub>). Evaporation of the mmol) and DMAP (286 mg, 2.34 mmol) was dissolved in and the residue was disolved in  $MeOH/EtoAc/H_2O$ . The materials) (400 mg, 1.17 mmol), H-(R,S)Itp(Ts) (See CH<sub>3</sub>CN (6 ml) and cooled to 5°C. EDC (236 mg, 1.23 Preparation of starting materials) (366 mg, 1.23 solvent resulted in a white solid, 688 mg (85%).

MS m/z 620 (M+ 1)

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(ii) H-(R)Cgl-Aze-(R,S)Itp(Ts)

product was dissolved in EtoAc/MeOH/ $\rm H_2^{2}O$  and the acidic solution was treated with 2 M NaOH(ag) to pH=8-9. The Evaporation of the solvent afforded 425 mg (100%) of solvent was removed by evaporation and the resulting through the solution for ca 4 min. After 45 min the Boc-(R)Cgl-Aze-(R,S)Itp(Ts) (500 mg, 0.8 mmol) was dissolved in  $\mathrm{CH}_2\mathrm{Cl}_2$  (50 ml) and HCl(g) was bubbled organic layer was separated and dried  $(\mathrm{Na}_2\mathrm{SO}_4)$  . the title compound as a white solid 2

MS m/z 520 (M+ 1)

25

(iii) BnOOC-CH2-(R)Cgl-Aze-(R,S)Itp(Ts)

CH<sub>3</sub>CN (5 ml) at 45°C. After a few hours the conversion increased to 60°C and an additional amount of Benzyl-Benzyl-2-(para-nitrobenzenesulfonyloxy)acetate (See 2-(para-nitrobenzenesulfonyloxy)acetate was added. mmol) and  $K_2CO_3$  (235 mg, 1.7 mmol) was stirred in Preparation of starting materials) (325 mg, 0.92 was only 25% and therefore the temperature was H-(R)Cgl-Aze-(R,S)Itp(Ts) (400 mg, 0.77 mmol), 35 3

was purified by RPLC. This gave 34 mg (7%) of the back-extraction of the acidic  $KHSO_4$ , some 340 mg which title compound  $\mathrm{KHSO_4}$ ,  $\mathrm{H_2O}$  and dried  $\mathrm{Na_2SO_4}$ ). This aforded, after combined organic phase was washed with  ${
m K_2CO_3}({
m sat})$ , 2 M phase was washed twice with EtOAc and then the residue. The phases were separated and the watersolvent was evaporated and  ${ t EtOAc/H_2O}$  was added to the The reaction was stirred for 48 h, (startingm.:product/25:63), and then worked up. The

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MS m/z 668 (M+ + 1).

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(iv)  $HOOC-CH_2-(R)CGl-Aze-(R,S)$  Itp x 2 HCl

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freeze-drying with 2.2 eq of 1 M HCl: contained the desired compound, 3 mg (10%) after which were analyzed with FAB-MS. Two fractions drying and preparative RPLC gave several fractions To the residue  $\rm H_2O$  and HOAc was added to pH=7.Freezewas removed and the  $\mathrm{NH_3}(1)$  was allowed to evaporate. appeared. The reaction was stirred for 5 min before it was quenched with HOAc (50  $\mu$ l). The dry-ice cooler cooler. Na(s) was added and a deep blue color was dissolved in THF (5 ml) and  $\mathrm{NH_3}(g)$  was destilled (40 ml) into the reaction flask with a dry-ice BnOOC-CH<sub>2</sub>-(R)Cgl-Aze-(R,S)Itp(Ts) (34 mg, 0.05 mmol)

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MS m/z 424 (M++1).

30

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Example 71

HOOC-CH2-(R) Cha-Aze-(R, 8) Itp

(i) Boc-(R)Cha-Aze-(R,S)Itp(Ts)

**3**5

Boc-(R)Cha-Aze-OH (See Preparation of starting

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gave 180 mg (57%) of the title compound gave 260 mg of crude product. Purification by RPLC Aze-(R,S)Itp(Ts) (See Example 70) case above. This night and worked up as described in the Boc-(R)Cglnigth. The reaction mixture was stirred an additional at room temperature over night. Extra (0.5 eq) Hmmol) was added and the resulting mixture was stirred (R,S) Itp(Ts) and EDC was added after stirring over mmol), DMAP (122 mg, 1 mmol) was dissolved in CH<sub>3</sub>CN (2.5 ml) and cooled to 5°C. EDC  $\times$  HCl (115 mg, 0.6 Preparation of starting materials) (155 mg, 0.52 materials) (169 mg, 0.5 mmol), H-(R,S)Itp(Ts) (See

MS m/z 634 (M++1).

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(ii) H-(R)Cha-Aze-(R,S)Itp(Ts)

25 20 Yield 163 mg (ca 100%): organic phase was dried( $\mathrm{Na_2SO_4}$ ) and evaporated to NaOH to pH=8-9. The phases were separated and the product was dissolved in  $\mathrm{CH_2Cl_2}$  and washed with 2 M solvent was removed by evaporation and the resulting through the solution for ca 4 min. After 45 min the dissolved in  $\mathrm{CH_2Cl_2}$  (20 ml) and  $\mathrm{HCl}(g)$  was bubbled Boc-(R)Cha-Aze-(R,S)Itp(Ts) (180 mg, 0.28 mmol) was

MS m/z 534 (M+ +1).

(iii) BnOOC-CH2-(R)Cha-Aze-(R,S)Itp(Ts)

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 $(Na_2SO_4)$ . Evaporation of the solvent gave a 171 mg of phase was washed with 10% citric acid and dried in EtOAc/ $\mathrm{H}_2\mathrm{O}$ . The phases were separated and organic solvent was evaporated and the residue was dissolved was stirred in CH<sub>3</sub>CN (1.5 ml) at 60°C for 2.5 h. The (45 mg, 0.33 mmol) and Br-CH<sub>2</sub>COOBn (39 mg, 0.17 mmol) H-(R)Cha-Aze-(R,S)Itp(Ts) (80 mg, 0.15 mmol),  $K_2CO_3$ 

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crude product, which was purified by RPLC yielding 53 mg (52%) of the title compound.

MS m/z 681 (M+ +1).

(iv) HOOC-CH2-(R)Cha-Aze-(R,S)Itp

Bnooc-CH<sub>2</sub>-(R)Cha-Aze-(R,S)Itp(TS) (50 mg, 0.07 mmol) was treated as described for Bnooc-CH<sub>2</sub>-(R)Cgl-Aze-(R,S)Itp(TS) (See Example 70 (iv)). This gave a product mixture which was purified on a RPLC yielding 12 mg of a 1:1 mixture of the title compound together with a reduced form which appear at mass 439 (m/z).

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15 MS m/z 438 (M\* +1)

Example 72

H-(R) Cha-Pic-(R,8) Itp x 2 ECl

(1) Boc-(R)Cha-Pic-(R,S)Itp(Ts)

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At roomtemperature 2.1 g (5.5 mmol) Boc-(R)Cha-Pic-OH (See Preparation of starting materials), 1.0 g (8.2 mmol) DNAP and 1.7 g (5.8 mmol) H-(R,S)Itp(TS) (See Preparation of starting materials) was dissolved in 40 mL acetonitrile. After a few minutes of stirring 1.1 g (5.8 mmol) EDC was added and the stirring was continued for 60 hours. The solvent was removed in vacuo and the residue was dissolved in CH<sub>2</sub>Cl<sub>2</sub>, washed with water, 0.3M KHSO<sub>4</sub> and KHCO<sub>3</sub> (aq) and dried(Na<sub>2</sub>SO<sub>4</sub>). Evaporation of the solvent and filtration through Silica gel gave 2.43 g (67%) of the product.

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MS m/z 661 (M+1)

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(ii) Boc-(R)Cha-Pic-(R,S)Itp

2.4 g (3.6 mmol) Boc-(R)Cha-Pic-(R,S)Itp(Ts) was dissolved in 15 mL THF and NH<sub>3</sub> (9) was condensed into the flask followed by addition of Na. The reaction was quenched after 5 min with acetic acid and the NH<sub>3</sub> and the THF was evaporated. The residue was freezedried from water and purified by RPLC

10 desired product.

the

 $(CH_3CN/0.1M\ NH_4OAC,\ 6/4)$  to give 0.93 g (51%) of

(iii) H-(R)Cha-Pic-(R,S)Itp x 2 HCl

13

MS m/z 507 (M +1)

At roomtemperature 50 mg (0.099 mmol) Boc-(R)Cha-Pic-(R,S)Itp was dissolved in ethylacetate saturated with HCl

(g). After stirring 2 h the solvent was removed in vacuo. The residue was freezedried from water three times to give 35 mg (74%) of the desired product.

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MS m/z 407 (M\* +1)

25 Example 73

HOOC-CH2-(R)Cha-Pic-(R,S)Itp x 2 HCl

(i) Boc-(R) Cha-Pic-(R,S) Itp(Z)

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At roomtemperature 0.84 g (1.66 mmol) Boc-(R)Cha-Pic-(R,S)Itp (See Example 72) was dissolved in 10 mL CH<sub>2</sub>Cl<sub>2</sub> and 10 mL 0.5M NaOH. 0.29 mL (1.82 mmol) 2-Cl was added dropwise. After stirring for 3 h the phases was separated and the organic phase was washed with water and dried over Na<sub>2</sub>SO<sub>4</sub>. Evaporation and flash chromatography (ethylacetate/heptane 9/1) gave 0.5 g

(47%) of the desired product.

MS m/z 641 (M++1)

### (ii) H-(R) Cha-Pic-(R, S) Itp(Z)

At roomtemperature 0.5 g (0.78 mmol) Boc-(R)Cha-Pic-(R,S)Itp(2) was dissolved in ethylacetate saturated with HCl. Water was added and the mixture was made basic with K<sub>2</sub>CO<sub>3</sub>. The phasees was separated. The waterphase was extracted with CH<sub>2</sub>Cl<sub>2</sub> and the organic phase was washed with water. The combined organic phase was then dried(Na<sub>2</sub>SO<sub>4</sub>). Evaporation of the solvent gave 0.3 g (71%) of the desired product.

MS m/z 541 (M++1)

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## (iii) Bnooc-CH<sub>2</sub>-(R)Cha-Pic-(R,S)Itp(Z)

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0.29 g (0.5 mmol) H-(R)Cha-Pic-(R,S)Itp(Z), 0.15 g (1 mmol) K<sub>2</sub>CO<sub>3</sub> was taken up in 25 mL acetonitrile. 154 mg (0.6 mmol) benzylbromoacetate was added and the mixture was stirred at 50°C for 4 h. evaporation and purification by RPLC

(acetonitrile:0.1M  $\mathrm{NH_4OAc}$  70:30) gave about 200 mg of the desired product.

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## (iv) $HOOC-CH_2-(R)Cha-Pic-(R,S)Itp x 2 HC1$

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200 mg BnOoC-CH<sub>2</sub>-(R)Cha-Pic-(R,S)Itp(Z) was dissolved in ethanol. A small spoon of 10% Pd on charcoal was added and the mixture was hydrogenated for 4 h. Filtration through hyflo, evaporation of the solvent followed by freezedrying from water gave 53 mg of the desired product.

**35** 

 $^{1}$ H NMR (300.13 MHz,  $D_{2}$ O); & 1.0-2.35 (overlapping m,

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<sup>22H</sup>), 3.28-3.51 (m, 5H), 3.51-3.64 (m, 1H), 3.75-4.03 (m, 3H), 5.03-5.14 (s broad, 1H). The signal of one of the protons is partially obscured by the H-O-D-signal.

MS m/z 465 (M+ +1)

Example 74

## H-(R)Cgl-Pro-(R,8)Hig x 2 HCl

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### (1) Boc-(R)Cgl-Pro-(R,S)Hig(Z)

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as eluent to yielded 1.1 g (59%) of the title stepwise gradient of  $ext{CH}_2 ext{Cl}_2/ ext{MeOH}$  (97/3, 95/5, 90/10 ) model 7924T ) using a 2mm silica plate with a crude product on a chromatotron (Harrison research, phase from above. Evaporation and purification of the dried ( $Na_2SO_4$ ), filtered and combined with the EtOAc then extracted with  $\mathrm{CH_2Cl_2}$ . The organic layer was  $(\mathrm{Na_2SO_4})$  and filtered. The oil and the water layer was organic layer. The ethyl acetate layer was dried a 0.3 M KHSO4-solution an oil separated from the was evaporated and the residue was dissolved in ethyl acetate. When the organic layer was washed twice with stirred at room temperature over night. The solvent added 0.62 g (3.2 mmole) of EDC and the mixture was Preparation of startingmaterials) in 15 ml  $\mathrm{CH_{2}Cl_{2}}$  was mmole) DMAP, 1.12 g (3.25 mmole) H- $\{R,S\}$ Hig(Z) (See (See Preparation of startingmaterials), 1.44 g (11.8 To a mixture of 1.0 g (2.95 mmole) Boc-(R)Cgl-Pro-OH

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## (ii) H-(R)Cgl-Pro-(R,S)Hig x 2 HCl

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81 mg (0.13 mmole) of Boc-(R)Cgl-Pro-(R,S)Hig(Z) was dissolved in 50 ml ethyl acetate saturated with HCl.

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solvent the residue was dissolved in  ${
m CH}_2{
m Cl}_2$  and washed layer was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and evaporated and mixture was stirred at 40°C for one hour and at room the crude product was purified on a chromatotron once with water and once with brine. The organic temperature over night. After evaporation of the bensylbromoacetate was mixed in 12 ml THF. The

95/5, 90/10 ) as eluent to yield 0.165 g (29%) of the (Harrison research, model 7924T ) using a 2mm silica plate with a stepwise gradient of  ${
m CH_2Cl_2/MeOH}$  (97/3, title compound. 2

(iii)  $HOOC-CH_2-(R)Cg1-Pro-(R,S)Hig \times .2 HC1$ 

evaporation of the solvent followed by freeze drying (R,S)Hig(Z) was mixed with 0.050 g Pd/C (5%), 0.7 ml hydrogenated at atmospheric pressure for four hours. 1 M HCl-solution and 10 ml ethanol. The mixture was Filtration of the catalyst through cellite and 0.165 g (0.25 mmole) of Bnooc-CH2-(R)Cgl-Pro-20 15

twice from water gave 0.1 g (75%) of the product.

 $^{1}$ H-NMR (D<sub>2</sub>O, 300 MHz): 6 1.05-1.45 (m, 5H), 1.55-2.5 (m, 15H), 3.08 (bt, 1H), 3.2-4.05 (m, 9H), 4.30 (d,

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 $^{13}\text{C-NMR}$  ( $^{}_{02}\text{O}$ , 75 MHz): carbonyl and guanidinecarbons: 6 154.9, 167.2, 169.4, 174.1 1H), 4.44 (m, 1H)

Example 76 30

H-(R)Cha-Pro-(R,8)Hig x.2 HCl

(i) Boc-(R)Cha-Pro-(R,S)Hig(Z)

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Preparation of starting materials), 0.95 g (7.8 0.72 g (1.95 mmole) Boc-(R)Cha-Pro-OH (See

H-NMR (D20, 300 MHz): 6 0.95-1.35 (m, 5H), 1.50-2.45 (m, 15H), 3.02 (bt, 1H), 3.1-3.8 (m, 7H), 4.13 (d, 3 times from water gave the title compound in 75% yield.

2

evaporation of the solvent followed by freeze drying

ethanol. 40 mg Pd/C (5%), 1 ml water and 0.5 ml 1 M

hydrogenated at atmospheric pressure over night. Filtration of the catalyst through cellite and

HC1-solution was added and the mixture was

evaporated and the residue was dissolved in 10 ml

The mixture was allowed to stand for one hour,

 $^{13}\text{C-NMR}$  (D20, 75 MHz): carbonyl and guanidinecarbons: 1H), 4.38 (bd, 1H)

\$ 154.8, 168.9, 174.4 12

MS m/z 393 (M+1)

Example 75 20

HOOC-CH2-(R) CG1-Pro-(R, B) Hig x 2 HC1

(1) H-(R)Cgl-Pro-(R,S)Hig(Z)

Example 74 (1)) was dissolved in 100 ml ethyl acetate the residue was dissolved in CH2Cl2. The organic layer saturated with HCl, and the mixture was allowed to stand for one hour. The mixture was evaporated and  $(\mathrm{Na_2SO_4})$  , filtered and evaporated to yield 0.825 g was washed twice with 0.2 M NaOH-solution, dried 1 g (1.6 mmole) Boc-(R)Cgl-Pro-(R,S)Hig(Z) (See (98%) of title compound. 30 25

(ii) BnOOC-CH2-(R)Cgl-Pro-(R,S)Hig(Z)

0.256 g (1.85 mmole)  $m K_2CO_3$  and 145  $\mu l$  (0.521 mmole) of

0.442 g (0.839 mmole) H-(R)Cgl-Pro-(R,S)Hig(Z), 32

mmole) DMAP, 0.74 g (2.14 mmole) 82% pure H-(R,S)Hig(Z) (See Preparation of starting materials) in 10 ml CH<sub>2</sub>Cl<sub>2</sub> was added 0.486 g (2.54 mmole) of EDC and the mixture was stirred at room temperature for 3 days. The mixture was diluted with CH<sub>2</sub>Cl<sub>2</sub> and washed with water, twice with 0.3M KHSO<sub>4</sub>-solution and once with brine. The organic layer was dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and evaporated and the crude product was purified by flash chromatography using CH<sub>2</sub>Cl<sub>2</sub>/MeOH 95/5 as eluent to yield 0.450 g (33%) of the product.

## (ii) H-(R)Cha-Pro-(R,S)Hig $\times$ 2 HCl

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50 mg (0.078 mmole) of Boc-(R)Cha-Pro-(R,S)Hig(Z) was dissolved in 20 ml ethyl acetate saturated with HCl. The mixture was allowed to stand for one hour, evaporated and the residue was dissolved in 10 ml ethanol. 20 mg Pd/C (5%) and 0.3 ml 1 M HCl-solution was added and the mixture was hydrogenated at atmospheric pressure for two hours. Filtration of the catalyst through through cellite and evaporation of the solvent followed by freeze drying twice from water gave 28 mg (76%) of the title compound.

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25 · 1H-NMR (D<sub>2</sub>O, 300 MHz): & 0.9-1.6 (m, 6H), 1.6-2.5 (m, 16H), 3.09 (t, 1H), 3.31 (t, 1H), 3.37-3.74 (m, 4H), 3.81 (m, 1H), 4.35-4.47 (m, 2H)

Example 77

H-(R)Cgl-Aze-Rig x 2 HCl

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(i) Boc-(R)Cgl-Aze-Rig(Z)

 $^{13}\text{C-NMR}$  (D<sub>2</sub>O, 75 MHz): carbonyl and guanidinecarbons: & 154.9, 169.8, 174.5

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chloride/methanol 9/1 to give 0.78 g (76%) of the desired compound after evaporation. through a pad of silica gel with methylene evaporated. The crude material was suction filtered sodium bicarbonate and water, dried  $(\mathrm{Na_2SO_4})$  and The methylene chloride layer was washed with aqueous potassium hydrogen sulfate and methylene chloride. days then evaporated and partitioned between aqueous hydrochloride. The reaction was allowed to stir for 3 of N-(3-dimethylaminopropyl)-N'-ethylcarbodiimide ml of dimethylformamide was added 0.33 g (1.7 mmol) dimethylaminopyridine in 30 ml of acetonitrile and 5 starting materials), 0.84 g (6.9 mmol) of mmol) of Boc-(R)Cha-Aze-OH( See preparation of Preparation of starting materials) , 0.59  $\mathbf{g}$  (1.6 To a solution of 0.50 g (1.6 mmol) of H-Rig(Z) (See

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<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>): 6 0.8-1.9 (m, 27 H), 2.4-2.6 (m, 2 H), 2.78 (bt, 2 H), 3.15-3.4 (m, 2 H), 3.80 (bt, 1 H), 4.0-4.4 (m, 4 H), 4.75 (bt, 1 H), 4.97 (bd, 1 H), 5.08 (s, 2 H), 7.1-7.4 (m, 7 H), 7.74 (b, 1 H).

## (ii) H-(R)Cgl-Aze-Rig(Z) x 2 HCl

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A flask containing Boc-(R)Cgl-Aze-Rig(Z), 0.76 g (1.2 mmol), in 50 ml of ethyl acetate was cooled in an ice bath. Dry HCl was bubbled through for 5 min and the solution was evaporated to give 0.74 g (100%) of the dihydrochloride as a white powder.

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1H-NMR (300 MHz, MeOD): & 1.1-2.0 (m, 18 H), 2.23 (m, 1 H), 2.68 (m, 1 H), 3.15-3.45 (m, 4 H), 3.72 (bd, 1 H), 3.9-4.0 (bd, 2 H), 4.27 (m, 1 H), 4.39 (m, 1 H), 4.78 (m, 1 H), 5.30 (s, 2 H), 7.3-7.5 (m, 5 H).

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(iii) H-(R)Cgl-Aze-Rig x 2 HCl

<sup>1</sup>H-NNR (300 MHz, D<sub>2</sub>0): 6 1.1-2.0 (m, 18 H), 2.37 (m, 1 H), 2.75 (m, 1 H), 3.08 (bt, 2 H), 3.39 (bt, 2 H), 3.8-4.0 (m, 3 H), 4.35-4.5 (m, 2 H), 4.90 (m, 1 H).

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 $^{13}\text{C-NMR}$  (75.5 MHz,  $^{12}\text{O}_2\text{O}$ ): guanidine and carbonyl carbons: 6 172.2, 169.4, 156.4.

#### 15 Example 78

## HOOC-CH2-(R) Cgl-Are-Rig x 2 HCl

## (i) BnOOC-CH<sub>2</sub>-(R)Cgl-Aze-Rig(Z)

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A mixture of 0.20 g (0.33 mmol) of H-(R)Cgl-Aze-Rig(2) (See Example 77), 0.13 g of potassium carbonate, 80 mg of sodium iodide, 10 ml of tetrahydro-furane and 10 ml of acetonitrile was heated at 60°C for 10 h. The solvents were evaporated and the crude material was flash chromatographed on silica gel using methylene chloride/methanol 92/8 as eluent. Yield: 0.13 g (58%).

30 lH-NMR (300 MHz, CDCl<sub>3</sub>) 6 0.9-2.1 (m, 18 H), 2.45 (m, 1 H), 2.61 (m, 1 H), 2.81 (m, 2 H), 2.88 (d, 1 H), 3.2-3.5 (m, 4 H), 3.94 (m, 1 H), 4.0-4.25 (m, 3 H), 4.85 (m, 1 H), 5.12 (s, 2 H), 5.14 (s, 2 H), 6.9-7.2 (b. 2 H), 7.2-7.5 (m, 10 H), 7.95 (m, 1 H).

32

(11) HOOC-CH<sub>2</sub>-(R)Cgl-Aze-Rig x 2 HCl

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A mixture of 0.12 g (0.18 mmol) of Bnooc-CH<sub>2</sub>-(R)Cgl-Aze-Rig(2), 5 ml of ethanol, 3 drops of conc. HCl and a small amount of 5% Pd/C was hydrogenated at atmospheric pressure for 1 h. The mixture was filtered through celite and evaporated. The residue was lyophilized in water to give 91 mg (98%) of the product.

10 H), 2.29 (m, 1 H), 2.70 (m, 1 H), 3.10 (m, 2 H), 3.34 (t, 2 H), 3.89 (dd, 2 H), 3.34 (t, 2 H), 3.83 (bd, 2 H), 3.89 (dd, 2 H), 4.00 (d, 1 H), 4.35 (m, 2 H), 4.87 (m, 1 H).

13C NMR (125.8 MHz, D<sub>2</sub>0): guanidine and carbonyl carbons: 6 171.8, 169.6, 167.7, 156.3.

15

#### Example 79

## HOOC-CH2-(R) Cha-Pro-Rig x 2 HC1

(i) Boc-(R)Cha-Pro-Rig(Z)

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To a solution of 0.25 g (0.82 mmol) of 4-aminoethyl1-benzyloxycarbonylamidino piperidine(H-Rig(Z)), (See preparation of starting materials), 0.32 g (0.82 mmol) of Boc-(R)Cha-Pro-OH (See Preparation of starting materials), 0.40 g (3.3 mmol) of dimethylaminopyridine in 10 ml of acetonitrile and 2 ml of dimethylformamide was added 0.165 g (0.86 mmol) of N-(3-dimethylformamide was added 0.165 g (0.86 mmol) of N-(3-dimethylaminopropyl)-N'-ethylcarbodiimide hydrochloride. The reaction was allowed to stir for 3 days then evaporated and partitioned between aqueous potassium hydrogen sulfate and methylene chloride. The methylene chloride layer was washed with aqueous sodium bicarbonate and water, dried (Na<sub>2</sub>SO<sub>4</sub>) and evaporated. The NRR spectrum of the crude product was satisfactory and the product which contained some

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<sup>1</sup>H-NMR (500 MHz, CDCl<sub>3</sub>) & 0.8-2.2 (m, 32 H; thereof 1.41 (s, 9 H)), 2.34 (m, 1 H), 2.77 (bt, 2 H), 3.10 (m, 1 H), 3.29 (m, 1 H), 3.40 (m, 1 H), 3.83 (m, 1 H), 4.17 (m, 2 H), 4.30 (m, 1 H), 4.54 (m, 1 H), 5.07 (m, 1 H), 5.08 (s, 2 H), 7.03 (m, 1 H), 7.05-7.4 (m, 7 H).

### (ii) H-(R)Cha-Pro-Rig(Z)

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A flask containing the crude product of Boc-(R)ChaPro-Rig(Z) in 100 ml of ethyl acetate was cooled in
an ice bath. Dry HCl was bubbled through for 5 min
and the solution was evaporated to get rid of the
excess of HCl. The product was dissolved in water and
the extracted twice with ethyl acetate to remove the
aqueous phase was made alkaline with NaHCO3 (aq) and
extracted twice with methylene chloride. The combined
organic phase was washed with water, dried (Na<sub>2</sub>SO<sub>4</sub>)
and evaporated. Yield: 0.37 g (81%) over two steps.

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## (iii) BnOOC-CH<sub>2</sub>-(R)Cha-Pro-Rig(Z)

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A mixture of 0.18 g (0.32 mmol) of H-(R)Cha-Pro-Rig(Z), an excess of potassium carbonate and 10 ml of acetonitrile was heated at 60°C for 2 h. The solvents were evaporated and the crude material was flash chromatographed on silica gel using methylene chloride/methanol 95/5 as eluent. Yield: 0.20 g

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(888).

<sup>1</sup>H-NMR (300 MHz, CDCl<sub>3</sub>) & 0.8-2.1 (m, 23 H), 2.37 (m, 1 H), 3.1-3.5 (m, 7 H), 4.0-4.2 (m, 2 H), 4.54 (m, 1 H), 5.1 (m, 4 H), 6.9-7.5 (m, 13 H).

(iv) Hooc-CH<sub>2</sub>-(R)Cha-Pro-Rig x 2 HCl

A mixture of 0.15 g (0.21 mmol) of BnOOC-CH2-(R)Cha10 Pro-Rig(Z), 10 ml of ethanol, 4 drops of conc. HCl
and a small amount of 5% Pd/C was hydrogenated at
atmospheric pressure for 1 h. The mixture was
filtered through celite and evaporated. The residue
was lyophilized in water to give 95 mg (64%) of the
product.

1H-NMR (500 MHz, MeOD) 6 0.85- 2.1 (m, 23 H), 2.30 (m, 1 H), 3.10 (m, 2 H), 3.25 (m, 1 H), 3.35 (m, 1 H), 3.54 (m, 1 H), 3.85-4.0 (m, 3 H), 4.03 (d, 1 H), 4.41 (m, 1 H), 4.50 (m, 1 H).

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<sup>13</sup>C-NMR (125.8 MHz, D<sub>2</sub>O): guanidine and carbonyl carbons: £ 174.0, 168.9, 168.1, 157.5.

#### 25 Example 80

HOOC-CH2-CH2-(R)Cha-Aze-Rig x 2 HC1

(i) Boc-(R)Cha-Aze-Rig(Z)

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To a solution of 0.25 g (0.82 mmol) of 4-aminoethyl-1-benzyloxy-cabonylamidino piperidine (H-Rig(2)), (See preparation of starting materials), 0.31 g (0.86 mmol) of Boc-(R)Cha-Aze-OH (See preparation of starting materials), 0.40 g (3.3 mmol) of dimethylaminopyridine in 10 ml of acetonitrile and 2 ml of dimethylformamide was added 0.17 g (0.86 mmol)

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(iii) Bnooc-CH2-CH2-(R)Cha-Aze-Rig(Z)

temperature for one week. It was evaporated and flash chloride/methanol 94/6 as eluent. Yield: 0.20 g chromatographed on silica gel using methylene ml of ethanol was allowed to stand at room

(m, 3 H), 4.84 (dd, 1 H), 5.05-5.15 (m, 4 H), 7.0-7.5 3.25 (m, 1 H), 3.31 (m, 1 H), 4.04 (q, 1 H), 4.1-4.2 (m, 18 H), 2.48 (m, 1 H), 2.54 (bt, 2 H), 2.68 (m, 2 <sup>1</sup>H NPR (500 MHz, CDCl<sub>3</sub>) 6 0.8-1.0 (m, 2 H), 1.1-1.8 H), 2.81 (bt, 2 H), 2.87 (m, 1 H), 3.20 (m, 1 H), 12

(iv) HOOC-CH2-CH2-(R)Cha-Aze-Rig x 2 HCl

The title compound was made and purified in the same mmol) of Bnooc-CH2-CH2-(R) Cha-Aze-Rig-(Z). Yield: 30 way as described in Example 80 from 0.20 g (0.28 mg (19%) of the dihydrochloride salt. 20

4.14 (t, 1 H), 4.17 (m, rotamer), 4.31 (m, 1 H), 4.46 3.3-3.4 (m, 4 H), 3.85 (bd, 2 H), 3.92 (m, rotamer), 1 H), 2.70 (m, 1 H), 2.83 (m, 2 H), 3.10 (m, 2 H), (m, 1 H), 4.89 (m, 1 H), 5.18 (m, rotamer). 25

 $^{13}\mathrm{C}$  NMR (125.8 MHz,  $^{12}\mathrm{O})$  guanidine and carbonyl carbons: 6 175.4, 171.8, 168.8, 156.3.

Example 81

HOOC-CH2-(R) Cha-Pro-(8) Itp x 2 HC1

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H), 4.03 (q, 1 H), 4.08 (m, 1 H), 4.18 (m, 2 H), 4.29

(m, 1 H), 4.78 (m, 1 H), 4.97 (m, 1 H), 5.09 (s, 2

H), 7.1-7.4 (m, 7 H), 7.65 (m, 1 H).

(11) H-(R)Cha-Aze-Rig(Z)

A flask containing the crude product of Boc-(R)Cha-20

excess of HCl. The product was dissolved in water and Aze-Rig(Z) in 100 ml of ethyl acetate was cooled in an ice bath. Dry HCl was bubbled through for 5 min and the solution was evaporated to get rid of the

phase was made alkaline with NaHCO3 (ag) and extracted dimethylformamide from the previous step. The aqueous the extracted twice with ethyl acetate to remove the twice with methylene chloride. The combined organic phase was washed with water, dried  $(\mathrm{Na}_2\mathrm{SO}_4)$  and evaporated. Yield: 0.31 g (70%) over two steps. 30 25

3.35 (m, 2 H), 4.05 (g, 1 H), 4.1-4.25 (m, 3 H), 4.86 'H-NMR (300 MHz, CDCl<sub>3</sub>) & 0.8-1.9 (m, 20 H), 2.48 (m, 1 H), 2.73 (m, 1 H), 2.85 (bt, 2 H), 3.25 (m, 1 H), (m, 1 H), 5.12 (s, 2 H), 6.9-7.2 (m, 2 H), 7.2-7.45 (m, 5 H), 7.93 (m, 1 H).

further purification.

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(bd, 1 H), 2.53 (m, 2 H), 2.77 (bt, 2 H), 3.25 (m, 2

H), 1.1-1.75 (m, 26 H; thereof 1.41 (s, 9 H)), 1.82

H-NMR (500 MHz, CDCl<sub>3</sub>) & 0.85 (m, 1 H), 0.97 (m, 1

12

hydrochloride. The reaction was allowed to stir for 3

of N-(3-dimethylaminopropyl)-N'-ethylcarbodiimide

days then evaporated and partitioned between aqueous

The methylene chloride layer was washed with aqueous

potassium hydrogen sulfate and methylene chloride.

dimethylformamide was used in the next step without

evaporated. The crude product which contained some

sodium bicarbonate and water, dried (Na2SO4) and

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Rig(Z) and 93 mg (0.57 mmol) of benzyl acrylate in 5 A solution of 0.31 g (0.57 mmol) of H-(R)Cha-Aze-

(m, 12 H), 8.03 (m, 1 H).

<sup>1</sup>H-NMR (500 MHz, CDCl<sub>3</sub>) 6 1.0-1.9 (m, 20 H), 2.33 (m,

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### (i) Boc-(R)Cha-Pro-(S)Itp(Ts)

G

the next step without further purification. about 60%)) of the desired product. Which was used in Evaporation gave 1.74 g (> 100% yiled (purity of dissolved in  $\mathrm{CH_2Cl_2}$ , washed with water, citric acid 9 (3.07 mmol) EDC was added. After 18 hours the (10%), KHCO $_3$  (aq), water and dried with Na $_2$ SO $_4$ . solvent was removed in vacuo and the residue wasin 12 mL acetonitrile. After stirring 20 minutes 0.59 (See preparation of starting materials) was dissolved (4.72 mmol) DMAP and 0.70 g (2.36 mmol) H-(S)Itp(Ts) OH (See preparation of startingmaterials), 0.78 g At roomtemperature 0.87 g (2.36 mmol) Boc-(R)Cha-Pro-

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FAB-MS:  $m/z = 647 (M^+ + 1)$ 

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### (11) H-(R)Cha-Pro-(S)Itp(Ts)

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as described for Boc-(R)Cha-Pic-(R,S)Itp(Z) (See Example 72 (ii)) to give 0.75 g (81%) of the title The Boc-protecting group was removed in the same way

FAB-MS:  $m/z = 547 (M^+ + 1)$ 

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## (iii) Bn00C-CH2-(R)Cha-Pro-(S)Itp(Ts)

mg of the desired product. ethylacetate/methanol 95/5 as eluent gave about 530 of the solvent followed by flash chromatography using the mixture was stirred at 50°C for 2 h. Evaporation 0.39 g (1.65 mmol) benzylbromoacetate was added and 0.75 g (1.37 mmol) H-(R) Cha-Pro-(S) Itp(Ts), 0.38 g (2.74 mmol)  $K_2CO_3$  was taken up in 15 mL acetonitrile.

FAB-MS:  $m/z = 695 (M^+ + 1)$ 

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(iv)  $Hooc-CH_2-(R)Cha-Pro-(S)$  Itp x 2 Hcl

the desired product after freeze-drying from aqueous water and the crude product was purified by RPLC was evaporated. The residue was freeze dried from after 30 min with acetic acid and the  $\mathrm{NH}_3$  and the THF (acetonitrile/0.1M HOAc 15/85) gave 0.25 g (61%) of flask and Na (m) was added. The reaction was quenched dissolved in 15 mL THF.  $NH_3$  (g) was condensed into the 0.53 g (0.76 mmol) BnOOC-CH<sub>2</sub>-(R)Cha-Pro-(S)Itp(Ts) was

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15 2H), 4.33-4.48 (overlapping m, 2H). 20H), 2.22-2.35 (m, 1H), 3.2-3.36 (m, 4H), 3.44-3.62 (overlapping m, 2H), 3.7-3.8 (m, 1H), 3.87-3.99 (m, 1H-NMR (500.13 MHz, D<sub>2</sub>0); 6 0.9-2.09 (overlapping m,

20 guanidinecarbons: & 154.3, 168.1, 169.0 and 174.2  $^{13}\text{C-NMR}$  (500.13 MHz,  $D_2\text{O}$ ); carbonyl- and

Example 82

### H-(R)Cha-Pro-(R,8)Nig x 2 HCl

(i) Boc-(R)Cha-Pro-(R,S)Nig(Z)

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30 by flash chromatography using  ${
m cH_2Cl_2/MeoH}$  95/5 as evaporated and the crude product was purified twice organic layer was dried ( $Na_2SO_4$ ), filtered and with 0.3 M KHSO $_4$ -solution and once with brine. The was diluted with  $\mathrm{CH_2Cl_2}$  and washed with water, twice the mixture was stirred for four days. The mixture  $\mathrm{CH_2Cl_2}$  and 117 mg (0.61 mmole) of EDC was added and Preparation of starting materials) was mixed in 2 mlmmole) DMAP, 130 mg (0.471 mmole) H-(R,S)Nig(Z) (See preparation of starting materials), 229 mg (1.87 174 mg (0.471 mmole) Boc-(R)Cha-Pro-OH (See

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MS m/z 627 (M+1)

(ii) H-(R)Cha-Pro-(R,S)Nig x 2 HCl

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0.1 ml 1 M HCl-solution was added and the mixture was 10 mg (0.016 mmole) of Boc-(R)Cha-Pro-(R,S)Nig(Z) was thydrogenated at atmospheric pressure for one and a dissolved in 15 ml ethyl acetate saturated with HCl. dissolved in 6 ml ethanol and 8 mg 5% Pd/C (5%) and The mixture was allowed to stand for half an hour. evaporation of the solvent gave 4 mg of the title The mixture was evaporated and the residue was half hour. After filtration through hyflo and compound the product

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<sup>1</sup>H-NWR (300 MHz, D<sub>2</sub>0): 6 0.9-1.58 (m, 6H), 1.58-2.45 (m, 13H), 2.65 (m, 1H), 3.19 (m, 1H), 3.34 (d, 2H), 3.4-3.73 (m, 4H), 3.82 (m, 1H), 4.34-4.49 (m, 2H). 20

13C-NMR(75 MHz, D20): carbonyl and guanidinecarbons: \$ 155.1, 169.9 and 174.8.

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#### Example 83

H-(R) Pro-Phe-Pab z 2 EC1

(1) Boc-(R)Pro-Phe-Pab(Z) ဗ္ဗ

preparation of starting materials) dissolved in 1 ml (13.91 mmol) DMAP in 40 ml  $\mathrm{CH}_3\mathrm{CN}$  at room temperature To a mixture of 1.2 g (3.31 mmol) Boc-(R) Pro-Phe-OH (See preparation of starting materials) and 1.70 g was added 0.98 g (3.35 mmol) H-Pab(Z) (See

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 $x\ 30\ ml\ Na_2CO_3,\ 1\ x\ 30\ ml\ water\ and\ dried.$  Evaporation of the solvent followed by flash chromatography using washed with 1 x 30 ml water, 3 x 30 ml 0.3 M KHSO<sub>4</sub>, 1 cooled to - 18°C and 0.66 g (3.48 mmol) EDC was added DMF. After stirring for 2 h the reaction mixture was  $\mathrm{CH_2Cl_2/MeOH}$  (95/5) as eluent gave 0.691 g (38%) of temperature over night. The solvent was evaporated and the residue was dissolved in 100 ml EtOAc and portion wise and the reaction was left at room

(11) H-(R)Pro-Phe-Pab(Z)

the title compound.

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and the combined organic phase was washed with water, NaOH. The washing water was extracted with 1 imes 25 ml 0.673 g Boc-(R) Pro-Phe-Pab(Z) was dissolved in 30 ml EtOAc and the solution was saturated with HCl(g) for and the organic phase was washed with 2  $\times$  20 ml 2 M Etoac which was combined with the other Etoac-phase evaporated and 60 ml EtOAc was added to the residue a few minutes (a white solid precipitated out from dried and evaporated to give 560 mg (98%) of the the solution). The solvent and excess HCl was desired product. 20 15

<sup>2</sup>H-NMR (500 MHz, CDCl<sub>3</sub>): 6 1.5-1.74 (m, 3H), 1.98-2.05 3.2 (ABX-system centered at 3.1, 2H), 3.62 (dd, 1H), 4.3-4.45 (ABX-system centered at 4.37, 2H), 4.58 (q, (m, 1H), 2.78-2.85 (m, 1H), 2.90-2.96 (m, 1H), 3.0-1H), 5.22 (s, 2H), 6.96 (bt, 1H), 7.1-7.4 (m, 10H), 7.46 (d, 2H), 7.76 (d, 2H), 8.12 (d, 1H).

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(iii) H-(R)Pro-Phe-Pab x 2 HCl

200 mg H-(R)Pro-Phe-Pab(Z) was dissolved in 10 ml 95 hydrogenated over 5 % Pd/C at atmospheric pressure % EtOH and 2 ml of water and the mixture was 35

for 5 h. Filtration of the catalyst and addition of 1 ml 1 M HCl followed by evaporation and freeze drying from water gave the title compound in 88 % yield.

<sup>1</sup>H-NMR (500 MHz, CD<sub>3</sub>OD): & 1.51-1.59 (m, 1H), 1.69-1.80 (m, 1H), 1.87-1.97 (m, 1H), 2.19-2.29 (m, 1H), 2.90 (dd, 1H), 3.20-3.33 (m, 3H, partially hidden by the solvent peak), 4.27 (m, 1H), 4.43-4.54 (AB-system centered at 4.48, 2H), 4.75-4.81 (m, 1H), 4.87 (s, 2H), 7.2-7.3 (m, 5H), 7.45 (d, 2H), 7.75 (d, 2H).

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 $^{13}\text{C-NMR}$  (125 MHz,  $D_2\text{O}$ ): amidine and carbonyl carbons: 6 166.7; 170.1 and 173.4.

#### Example 84

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### HOOC-CH2-(R)Pro-Phe-Pab x 2 HC1

### (i) BnOOC-CH<sub>2</sub>-(R)Pro-Phe-Pab(Z)

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To a slurry of 244 mg (0.463 mmol) H-(R)Pro-Phe-Pab(Z) (See Example 83) and 159.9 mg (1.157 mmol) K<sub>2</sub>CO<sub>3</sub> in 8 ml DMF/CH<sub>3</sub>CN (5/3) was added 127.2 mg (0.555 mmol) benzylbromo accetate dissolved in 2 ml DMF and the mixture was stirred at 60°C for 1.5 h and at room temperature over night. The solvent was evaporated and the residue was dissolved in 50 ml EtOAc, washed with 2 x 20 ml water and dried (Na<sub>2</sub>SO<sub>4</sub>). Evaporation of the solvent followed by flash chromatography using CH<sub>2</sub>Cl<sub>2</sub>/MeOH (9/1) as eluent gave 176 mg (56 %) of the title compound as a white solid.

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1H-NMR (300 MHz), CDCl<sub>3</sub>): & 1.45-1.80 (m, 3H), 2.06 (m, 1H), 2.54 (m, 1H), 2.92-3.28 (m, 6H), 4.3-4.5 (ABX-system centered at & 4.4, 2H), 4.60 (dd, 1H), 5.10 (apparent s, 2H), 5.2 (apparent s, 2H), 7.1-7.4 (m, 15H), 7.43 (d, 2H), 7.75 (d, 2H), 7.932 (d, 1H).

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## (ii) HOOC-CH2-(R)Pro-Phe-Pab x 2HC1

170 mg (0.252 mmol) of BnOOC-CH<sub>2</sub>-(R)Pro-Phe-Pab(Z) was dissolved in 12 ml EtOH/water (5/1) and hydrogenerated over 5 % Pd/C at atmospheric for 4.5 h. The catalyst was filtered off, the solvent evaporated and the residue freeze dried from HCl(aq) to give the title compound.

<sup>13</sup>C-NMR (125 MHz, D<sub>2</sub>O): amidine and carbonyl carbons: δ 166.8, 169.1, 169.5 and 173.2.

#### Example 85

#### H-(R)Phe-Phe-Pab

### (i) Boc-(R)Phe-Phe-Pab(Z)

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Boc-(R)Phe-Phe-OH (16.4 mmol) (see preparation of starting materials), Pab(Z)-HCl (18.0 mmol) and 4-dimetylaminopyridine (24.6 mmol) were dissolved in 10e-water temperature and 1-(3-dimetylaminopropyl)-3-added. The cooling bath was removed and the reaction mixture was stirred over night. The solvent was then dissolved in 50 mL of ethylacetate and the residue solution extracted with 50 mL of water. Boc-(R)Phe-

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Phe-Pab(Z) precipitating from the two-phase mixture

**\*** 

12H), 4.64 (m, 1H), 4.44 (m, 2H), 4.13 (t, 1H), 3.1-2.8 (m, 4H).

Example 86

HOOC-CO-(R) Phe-Phe-Pab

(i) MeOOC-CO-(R) Phe-Phe-Pab(Z)

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H-(R)Phe-Phe-Pab(Z) (0.87 mmol) (see Example 85 (ii)) was dissolved in 10 mL of tetrahydrofuran. The solution was cooled on an icewater bath and triethylamine (1.73 mmol) followed by methyloxalylchloride (0.95 mmol) were added. The cooling bath was removed and the reaction mixture stirred for 18 h at ambient temperature. The reaction mixture was diluted with ethylacetate and extracted with water. The organic phase was collected and the solvent was removed under reduced pressure yielding 0.45 g of MeoOC-CO-(R)Phe-Phe-Pab(Z) (78%) which was used in the next step without further purification. TSP-MS found m/z 664 (calculated for MH\* (C37%38N5O))

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(ii) HOOC-CO-(R) Phe-Phe-Pab(Z)

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MeOOC-CO-(R)Phe-Phe-Pab(Z) (0.68 mmol) was dissolved in 4 mL of tetrahydrofuran and 2 mL of water. Lithiumhydroxide (2.6 mmol) was added and the reaction mixture was stirred at room temperature for 1.5 h. After complete hydrolysis the reaction mixture was diluted with 25 mL of water and acidified by addition of 0.5 mL of acetic acid. The precipitate was filtered and washed with several portions of water yielding 0.40 g of crude HOOC-CO-(R)Phe-Pab(Z) after drying under vacuum at 45°C for 24 h. The crude product was slurried in 10 mL of ethanol

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was filtered and washed with water yielding 8.7 g (78%) after drying under vacuum at 45°C for 24 h.  $^{1}$ H NMTR (200 MHz, d-CHCl<sub>3</sub> and d4-CH<sub>3</sub>OH);  $\delta$  8.35-7.00 (m, 19H), 4.63 (t. 1H), 4.3-4.1 (m, 1H), 3.40-2.70 (m, 6H), 1.30 (s, 9H).

(ii) H-(R)Phe-Phe-Pab(Z)

dissolved in a mixture of 50 mL of methylenechloride, Boc-(R) Phe-Phe-Pab(Z) (10.3 mmol) was slurried in 70 5.0 g of H-(R)Phe-Phe-Pab(Z) (84%). <sup>1</sup>H NMR (200 MHz, solvent was removed under reduced pressure yielding d<sub>6</sub>-DMSO); 6 9.1 (8, 2H), 8.59 (m, 1H), 8.1 (m, 1H), 7.90 (d, 2H), 7.4-7.0 (m, 17H), 5.09 (s, 2H), 4.58 (R) Phe-Phe-Pab(Z) was filtered off and washed with ethylacetate/HCl was added. The slurry was stirred ethanol. The organic layer was collected and the for 4 h after which the hydrochloride salt of Hserveral portions of ethylacetate. The salt was 50 mL of 1 M potassiumcarbonate and ca 5 mL of (m, 1H), 4.31 (m, 2H), 3.1-2.7 (m, 4H). ml of ethylacetate and 31 mL of 3.3 M 20 2 12

(80:20:2). Yield 76 mg of the title compound (41%).  $^{1}\mathrm{H}$ off. Evaporation of the solvents gave crude H-(R)Phehydrogen pressure in a Parr shaking apparatus for 2 days. After complete hydrogenolysis the mixture was diluted with methanol and the catalyst was filtered (iii) H-(R)Phe-Phe-Pab(Z) (0.42 mmol) was dissolved solution and the mixture was hydrogenated at 45 psi TATR (200 MHz, d<sub>6</sub>-DMSO); 6 7.61 (d, 2H), 7.4-7.0 (m, Palladium on charocoal (42 mg) was charged to the Phe-Pab which was purified by chromatography on in 10 mL of tetrahydrofuran and 1 mL of water. methylenechloride-methanol--ammoniumhydroxide neutral alumina (70-230 Mesh) eluting with ဗ္ဗ 35 25

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5.10 (s, 2H), 4.54 (m, 2H), 4,34 (m, 2H), 3.2-2.6 (m, 2H), 8.41 (d, 1H), 7.89 (d, 2H), 7.4-6.9 (m, 17H), over two steps).  $^{1}\text{H}$  NMR (200 MHz,  $d_{6}\text{-DMSO}$ );  $\delta$  8.62 (m, yielding 0.23 g of HOOC-CO-(R)Phe-Phe-Pab(Z) (41% and the insoluble title compound was filtered off, and 1 mL of water. The solution was brought to reflux

### (iii) HOOC-CO-(R)Phe-Phe-Pab

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7.35-6.8(m,12H), 4.6-4.0(m, 4H), 3.0-2.6(m, 4H). 6 9.2(s), 8.78(d), 8.60(m), 7.91(m), 7.79(d, 2H), title compound (49%). 1H NMR (200 MHz, d6-DMSO); off. Evaporation of the solvents yielded 50 mg of the with 40 mL of methanol and the catalyst was filtered After complete hydrogenolysis the mixture was diluted pressure in a Parr shaking apparatus for 2 days. the mixture was hydrogenated at 45 psi hydrogen on charcoal (52 mg) was charged to the solution and 20 mL of tetrahydrofuran and 5 mL of water. Palladium HOOC-CO-(R)Phe-Phe-Pab(Z) (0.20 mmol) was slurried in

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#### Example 87

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HOOC-CH2-(R) Phe-Phe-Pab

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### (i) BnOOC-CH2-(R)Phe-Phe-Pab(Z)

and from the collected organic phase the title solution was rapidly extracted with 10 mL of water the residue dissolved in 10 mL of ethylacetate. The After complete alkylation the solvent was removed and to 30°C and stirred at that temperature for 2 days. was added to the mixture and the solution was heated 10 mL of acetonitrile. Iodobenzylacetate (0.95 mmol) and potassium carbonate (2.6 mmol) were slurried in H-(R)Phe-Phe-Pab(Z) (0.87 mmol) (see Example 85 (ii))

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6.55(t, 1H), 5.21(s, 2H), 5.03(s, 2H), 4.64(m, 1H), NMR (200 MHz, CDCl<sub>3</sub>); & 7.79(d, 2H), 7.5-7.1(m, 22H), 4.41(m 2H), 3.3-2.6(m, 7H). Yielding 177 mg Bnooc-CH $_2$ -(R)Phe-Phe-Pab(Z) (28%).  $^1\mathrm{H}$ filtered off and dried under vacuum at 45°C for 24 h compound precipitates. BnOOC-CH2(R)Phe-Phe-Pab(2) was

## (ii) BnOOC-CH2-(R)Phe-Phe-Pab(Z)

20 5 10 mg of the title compound (59%). TSP-MS found m/z 502 filtered off. Evaporation of the solvents yielded 95 diluted with 40 mL of water and the catalyst was days. After complete hydrogenolysis the mixture was (calculated for MH $^{+}$  (C<sub>28</sub>H<sub>32</sub>N<sub>5</sub>O<sub>4</sub>)502). hydrogen pressure in a Parr shaking apparatus for 2 solution and the mixture was hydrogenated at 45 psi Palladium on charcoal (41 mg) was charged to the  $Bn00C-CH_2-(R)Phe-Phe-Pab(Z)$  (0.32 mmol) was slurried in 30 mL of tetrahydrofuran and 3 mL of water.

#### Example 88

#### H- (R) Cha-Pro-Mig

### Boc-(R)Cha-Pro-Mig(Z)

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evaporated. The crude product was purified by flash The organic layer was dried with  $Na_2SO_4$  and in EtOAc and washed with  ${
m H_2O}$ ,  ${
m NaHCO_3}$  (aq) and brine. The CH<sub>3</sub>CN was evaporated and the residue was dissolved allowed to reach roomtemperature and left for 5 days. mmol) of EDC at -10°C. The reaction mixture was mmol) of DMAP in 10 mL  $\mathrm{CH_{3}CN}$  was added 0.232 g (1.21) preparation of starting materials) and 0.227 g (1.86 materials), 0.245 g (0.93 mmol) of H-Mig(Z) (see (R)Cha-Pro-OH (see preparation of starting To a stirred mixture of 0.344 g (0.93 mmol) Boc-

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chromatography using a gradient of EtOAc/MeOH, 95/5 to 90/10, as eluent to yield 0.340 g (60 %) of the title compund.

(ii) H-(R)Cha-Pro-Mig(Z)

0.34 g (0.55 mmol) Boc-(R)Cha-Pro-Mig(Z) was dissolved in 8 mL of EtoAc saturated with Hcl(g) and stirred for 10 min. at roomtemperature. 10 mL of a saturated solution of KOH(aq) was added dropwise. The layers were separated and the aqueous phase was extracted with 3x8 mL EtoAc. The organic layers were combined, washed with brine, dried with Na<sub>2</sub>SO<sub>4</sub> and evaporated to yield 0.286 g (100 %) of the title

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(iii) H-(R)Cha-Pro-Mig

combound.

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0.050 g (0.132 mmol) of H-(R)Cha-Pro-Mig(Z) was dissolved in 3 mL MeOH and hydrogenated over 10 % pd/C at atmospheric pressure over night. The solution was filtered through celite and the solvent evaporated to yield 0.040 g (80 %) of the title compound.

20

JH-NMER (500 MHz, MeOD): 6 0.92-1.02 (m, 2H), 1.18-1.47 (m, 6H), 1.66-1.73 (m, 4H), 1.85-2.04 (m, 4H), 2.17-2.22 (m, 1H), 2.95-2.98 (m, 1H), 3.12-3.16 (m, 1H), 3.47-3.55 (m, 2H), 3.62-3.66 (m, 1H), 3.75-3.78 (m, 1H), 3.85-3.89 (m, 1H), 4.05-4.12 (m, 3H), 4.34-4.37 (m, 1H).

25

Signals from a minor rotamer appear at: 6 3.4, 3.7, 4.13-4.16, 4.3.

MS m/z 379 (M+ 1)

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H-(R)Cha-Pro-Dig

Example 89

5 (1) Boc-(R)Cha-Pro-Dig(Z)

To a stirred mixture of 0.280 g (0.76 mmol) Boc-(R)cha-Pro-OH (see preparation of starting materials), 0.210 g (0.76 mmol) of H-Dig(Z) (see preparation of starting materials) and 0.186 g (1.52 mmol) of DMAP in 8 mL CH<sub>3</sub>CN was added 0.189 g (0.99 mmol) of EDC at -10°C. The reaction mixture was allowed to reach roomtemperature and left for 4 days. The CH<sub>3</sub>CN was evaporated and the residue was dissolved in EtoAc and washed with H<sub>2</sub>O, NaHCO<sub>3</sub> (aq) and brine. The organic layer was dried with Na<sub>2</sub>SO<sub>4</sub> and evaporated. The crude product was purified by flash chromatography using a

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gradient of EtOAC/MeOH, 95/5 to 90/10, as eluent to yield 0.210 g (44 %) of the title compund.

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(11) H-(R) Cha-Pro-Dig(Z)

0.210 g (0.33 mmol) Boc-(R)Cha-Pro-Dig(2) was dissolved in 8 mL of EtOAc saturated with HCl(g) and stirred for 10 min. at roomtemperature. 8 mL of a saturated solution of KOH(ag) was added dropwise. The layers were separated and the aqueous phase was extracted with 3x8 mL EtOAc. The organic layers were combined, washed with brine, dried with Na<sub>2</sub>SO<sub>4</sub> and evaporated to yield 0.146 g (83 %) of the title compound.

(iii) H-(R)Cha-Pro-Dig

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0.046 g (0.087 mmol) of H-(R)Cha-Pro-Dig(Z) was dissolved in 3 mL MeOH and hydrogenated over 10 %

evaporated to yield 0.040 g (100 %) of the title was filtered through celite and the solvent Pd/C at atmospheric pressure over night. The solution

3H), 4.07-4.25 (m, 3H), 4.35-4.39 (m, 2H). 3.15-3.29 (m, 1H), 3.44-3.57 (m, 2H), 3.65-3.87 (m, 2.21 (m, 1H), 2.74-2.83 (m, 1H), 2.94-2.99 (m, 1H), (m, 6H), 1.66-1.74 (m, 4H), 1.78-2.05 (m, 4H), 2.13-1H-NMR (500 MHz, MeOD): & 0.90-1.04 (m, 2H), 1.10-1.47

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Signals from a minor rotamer appear at: & 4.29-4.32.

MS m/z 393 (M+ + 1)

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Example 90

H-(R)Cha-Aze-Dig

(i) Boc-(R)Cha-Aze-Dig(Z)

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Pro-Dig(2) in a yield of 0.253 g (54 \$). OH and H-Dig(Z) (see preparation of starting material) according to the procedure for Boc-(R)Cha-The title compound was prepared from Boc-(R)Cha-Aze-

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(ii) H-(R)Cha-Aze-Dig(Z)

Dig(2) in a yield of 0.210 g (100 %). Dig(Z) according the procedure for Boc-(R)Cha-Pro-The title compound was prepared from Boc-(R)Cha-Aze-

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(iii) H-(R)Cha-Aze-Dig

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Pd/C at atmospheric pressure over night. The solution dissolved in 3 mL MeOH and hydrogenated over 10 % 0.060 g (0.117 mmol) of H-(R)Cha-Aze-Dig(Z) was

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evaporated to yield 0.042 g (95 %) of the title was filtered through celite and the solvent compound.

3.39-3.44 (m, 1H), 3.72-3.80 (m, 2H), 4.01-4.04 (m, 1H), 4.14-4.23 (m, 2H), 4.48-4.49 (m, 1H), 4.60-4.64 2.68 (m, 1H), 2.80-2.83 (m, 1H), 3.14-3.29 (m, 1H), (m, 1H). (m, 6H), 1.66-1.90 (m, 8H), 2.15-2.17 (m, 1H), 2.66-<sup>1</sup>H-NWR (500 MHz, MeOD): £ 0.91-1.02 (m, 2H), 1.18-1.48

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Signals from a minor rotamer appear at: 6 2.25, 2.6, 4.3, 4.67.

15  $MS m/z 379 (M^+ + 1)$ 

Examples of pharmaceutical preparations

25 20 suspension for parenteral use. Liquid solid or semisolid dosage forms for topical administration. or modified release tablets. Liquid or solid-Lyophilized substance or liquids as emulsion or semisolid dosage forms for rectal administration. administration such as plain tablets, coated tablets formulated in solid dosage forms for oral The compound according to the invention can be

oral or masal inhalation. In pressurized aerosols or in dry powder inhalers for

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<u>Example P1</u>

Tablets for oral administration

ingredients: 1000 tablets are prepared from the following

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bunoamos exitor	100 9	
ACCIVE COMPANY	200 9	
nactors	30 9	
winnerstalline cellulose	30 ଫ	
Microsium Stearate	פי	

The active constituent and lactose are mixed with an aqueous solution of polyvinyl pyrrolidone. The mixture is dried and milled to form granules. The microcrystalline cellulose and then the magnesium stearate are then admixed. The mixture is then compressed in a tablet machine giving 1000 tablets, each containing 100 mg of active constituent.

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### Example P2

# Solution for parenteral administration

20 A solution is prepared from the following ingredients:

Active compound Sodium chloride for injection 6 Sodium hydroxide for pH adjustment ad pH 5-7 Water for inj. up to 1000 ml

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The active constituent and the sodium chloride are dissolved in the water. The pH is adjusted with 2 M NaOH to pH 3-9 and the solution is filled into sterile ampoules.

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Example P3

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<u> Tablets for oral administration</u>

150 g	20 g	120 g	20 g	S G	3 9
Active compound	Sodium aluminium silicate	Paraffin	Microcrystalline cellulose	Hydroxy propyl cellulose	Sodium stearyl fumarate
;	2	, ,	4	'n,	. 6
				ហ	

1-4 are mixed and an aqueous solution of 5 is added. The mixture is dried and milled and 6 is admixed. The mix is then compressed in a tablet machine.

### Example B6

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## Inhaler powder

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The active compound is micronized in a jet mill to a particle size suitable for inhalation (mass diameter  $< 4 \mu m$ ).

20 100 mg of the micronized powder is filled into a powder multidose inhaler (Turbohaler®). The inhaler is equipped with a dosing unit which delivers a dose of 1 mg.

### Biology

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# Determination of Thrombin clotting Time (TT):

Human thrombin (T 6769, Sigma Chem Co) in buffer solution, pH 7.4, 100 μl, and inhibitor solution, 100 μl, were incubated for one min. Pooled normal citrated human plasma, 100 μl, was then added and the clotting time measured in an automatic device (KC 10, Amelung).

The clotting time in seconds was plotted against the inhibitor concentration, and the  ${\rm IC}_{\rm SO}{\rm TT}$  was determined

by interpolation.

 $\mathrm{IC}_{50}\mathrm{TT}$  is the concentration of inhibitor that doubles the thrombin clotting time for human plasma.

Determination of Activated Partial Thromboplastin

APTT was determined in pooled normal human citrated plasma with the reagent PTT Automated 5 manufactured by Stago. The inhibitors were added to the plasma (10 determined in the mixture by use of the coagulation of the reagent producer. The clotting time in seconds plasma and the IC50APTT was determined against the inhibitor concentration in interpolation.

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20 IC<sub>50</sub>APTT is defined as the concentration of inhibitor in plasma that doubled the Activated Partial Thromboplastin Time.

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# <u>Determination of thrombin time ex vivo</u>

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The inhibition of thrombin after oral administration of the compounds were examined in conscious rats that two days prior to the experiment were equipped with a Catheter for blood sampling from the carotid artery. On the experimental day blood samples were withdrawn at fixed times after the administration of the compound into plastic tubes containing 1 part sodium citrate solution (0.13 mol per L.) and 9 parts of poor plasma. The plasma was used for determination of thrombin time as described below.

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The citrated rat plasma, 100  $\mu$ l, was diluted with a saline solution, 0.9%, 100  $\mu$ l, and plasma coagulation was started by the addition of human thrombin (T 6769, Sigma Chem Co, USA) in a buffer solution, pH 7.4, 100  $\mu$ l. The clotting time was measured in an automatic device (KC 10, Amelumg, Germany).

Determinaton of the inhibition constant  $\mathbf{K}_i$  for plasma kallikrein

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K<sub>1</sub> determinations were made with a chromogenic substrate method, and performed on a Cobas Bio centrifugal analyzer manufactured by Roche (Basel, Switzerland). Residual enzyme activity after incubation of human plasma kallikrein with various concentrations of test compound was determined at three different substrate concentrations, and measured as change in optical absorbance at 405 nm and 37°C.

15

Human plasma kallikrein (E.C.3.4.21.34, Chromogenix AB, Mölndal, Sweden), 250 µl of 0.4 nkat/ml in buffer (0.05 mol/1 Tris-HCl, pH 7.4, 1 0.15 adjusted with NaCl) with bovine albumin 5 g/l (cat no 810033, ICI Biochemicals Ltd, High Wycombe, Bucks, GB), was incubated for 300 s with 80 µl of test compound solution in 0.15 mol/1 NaCl containing albumin 10 g/l. An additional 10 µl of water was supplied in this step. Then 40 µl of kallikrein substrate (S-2302, Chromogenix AB, 1.25, 2.0 or 4.0 mmol/1 in water) was added together with another 20 µl of water, and the absorbance change monitored.

 $K_1$  was evaluated from Dixon plots, i.e. diagrams of inhibitor concentration versus 1/ ( $\Delta A/min$ ), where the data for the different substrate concentrations form straight lines which intercept at  $x=-K_1$ .

		PCT/SE94/00535 W	WO 94/29336		PCT/SE94/00535	
W0 5	WO 94/29336	223				
			Gly =		glycine	
	ARREVIATIONS		8		hours	
			HCl =		hydrochloric acid	
		acetyl	Hex =		hexyl	
	2 1	aqueous	HOAC =		acetíc acid	
,	ı fo	ne-2-carboxylic acid	HOBt =		N-hydroxy benzotriazole	
n N		piperidine-3-carboxylic acid	Hoc		Homocyclohexyl alanine	
	betaPic =	tout = hit v lox y carbony 1	n C		Homophenyl alanine	
		(n=+ert-butvloxycarbonyl-	don HOSh		N-hydroxysuccinimide	
	Boc-Dig(Z)	sminoethyl)-1-(N-benzyloxy-			3-aminoethyl-1-(N-benzyloxycarbonyl-	
		-			amidino) azetidine	
9	1 (6) 2 (7)	3-(N-ter-butyloxycarbonyl-	H-Dia a		3-aminoethyl-1-amidino azetidine	
	Boc-m19(4) =	aminomethv1)-1-(N-benzyloxy-	u_(2)Hig(2)=	=(2)	(3RS)-1-(N-benzyloxycarbonylamidino)-	
		carbonylamidino) azetidine	F== (2 (w) - 11		3-aminoethyl pyrrolidine	
			, a a) _ u		(3RS)-1-amidino-3-aminoethyl	
	Boc-Pig(Z) =	4-(N-tertument tox) of the part of the par			pyrrolidine	
15		aminomethyl)-1-(N-benzylowy			pyrrotrans	
ł		carbonylamidino) piperidine	H-Hig =		1-abidino-3-geninoeda y 171-0-17	
	Boc-Pig(Z), =	4-(N-tert-butyloxycarbonyl-	H-(R,S)Itp(Ts)=	p(Ts)=	(4RS)-1,3-diaza-z-tosylimino-4-	
	7/-)6-1-009	aminomethyl)-1-(N,N'-dibenzyloxy-			aminoethylcyclohexane	
		carbonylamidino) piperidine	0 H-(R,S)Itp	ij Ω.	(4RS)-1,3-diaza-2-imino-4-	
1		u,			aminoethylcyclohexane	
20	PLTHE -	henzyl	H-(S)Itp(Ts)=	T8)=	(4S)-1,3-diaza-2-tosylimino-4-	
					aminoethylcyclohexane	
	Bu a	Authority alveine	u=(S) Itp	U	(4S)-1,3-diaza-2-imino-4-aminoethyl-	
	cg] =	9	4 · · · · · ·		cyclohexane	
	cha =	p-cyclonexy argument (25			1 1-diaza-2-imino-4-aminoethyl	
25	CME-CDI =	1-Cyclonexy1-3-(2-morphosos)10nate	a dat-H	•	ryclohexane	
		Carroon in the metric of the m			- N - L - C - C - C - C - C - C - C - C - C	
	DBU =	1,8-diazabicyclolo.4.0jumec-/-cmc	H-M1g(Z)		J-aminomorphy - (" ) azetidine	
	a DCC	dicyclohexyl carbodilmide			Denzy Long car bond remains a set idine	
	DCII a		30 H-Mig =	-	3-abinometnyl-l-amidino acettariio	
,		N,N-dimethyl amino pyridine	H-(R,S)N1g(Z)=	1g(Z)=	(3RS)-1-(N-benzyloxycarbonylamiaring)	
2		dimethyl formamide			3-aminomethyl pyrrolldine	
	i jed	Almethyl sulphoxide	H-(R,S)N19	19 =	(3RS)-1-amidino-3-aminomethyl	
	DMSO =	-(3-pimetvlaminopropvl)-3-			pyrrolidine	
	EDC =	ride	# Civin		1-amidino-3-aminomethyl pyrrolidine	
					1-amidino-4-aminomethyl benzene	
35	Et n	ethyl	H-Pan a			
	EtOAc =	ethyl acetate				
	EtoH =	ethanol				

		225
	H-Pab(Z) =	4-aminomethyl-1-(N-benzyloxy
	H-Pac = H-Pac(Z) =	1-amidino-4-aminomethyl cyclohexane
Ú	H-Pig =	4-aminomethyl-1-(N-benzyloxy Carbonylamidino) cyclohexane 4-aminomethyl-1-amidino -income
	H-Pig(Z) =	4-aminometal :

 $H-Pig(Z)_2 =$ H-Pig(Z) =

amidino)-piperidine

4-aminomety1-1-(N-benzyloxycarbony1-

10

H-Rig(Z) = H-Rig =

MeOH =

4-aminoethyl-1-N-amidino piperidine

carbonylamidino)piperidine 4-aminoethyl-1-(N-benzyloxycarbonylamidino) piperidine 4-aminomethyl-1-(N,N'-dibenzyloxy

15

Pd/C = MA " XS G Мра =

mega pascal

methanol methy1

Pic = Pg1 = Phe =

> phenyl glycine palladium on charcoal N-methyl morpholine

20

RPLC = Pro =

> pipecolinic acid phenyl alanine

proline

25

Tf =

chromathography Reverse phase high performace liquid

trifluoroacetic acid trifluoromethylsulfonyl

tetrahydrofuran

THF = TFA =

11c =

1-carboxy-1,2,3,4-tetrahydroisoquinoline

tosyl

valine

benzyloxy carbonyl

2 0 Val = Ts =

35 Prefixes n, s, i and t have their usual meanings: normal, iso, sec and terriary. The stereochemistry for the amino acids is by default (S) if not otherwise stated.

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ABBREVIATIONS (continued, the wavy lines on the nitrogen atoms in the structural formulas below signify the bond position of the fragment.)

Mig (n=1) Dig (n=2)

Itp (n=2)

Nig (n=1) Hig (n=2)

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CLAIMB

1. A compound of the general formula

-M-(CH2)-B

Formula 1

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wherein:

A<sup>1</sup> represents a structural fragment of Formula IIa, IIb, IIc, IId or IIe; 15

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wherein:

k is an integer 0, 1, 2, 3 or 4;

m is an integer 1, 2, 3 or 4;

q is an integer 0, 1, 2, or 3;

carbon atoms, and  $R^{11}\ is\ H\ or\ an\ alkyl\ group\ having\ 1\ to$  ${\rm CONIR}^{12}$ , where  ${\rm R}^{12}$  is H or an alkyl group having 1 to 4 atoms, or  $R^{11}00C$ -alkyl-, where the alkyl group has 1 to alpha substituent is a group  $R^{17}-(CH_2)_{\mathfrak{p}^-}$ , wherein  $\mathfrak p$  is position which is alpha to the carbonyl group, and the  $R^{\rm l}$  represents H, an alkyl group having 1 to 4 carbon 4 carbon atoms and is possibly substituted in the 0, 1 or 2 and  $\mathbb{R}^{17}$  is methyl, phenyl, OH, COOR<sup>12</sup>, 6 carbon atoms, or

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 $\rm R^1$  represents Ph(4-COOR  $^{12}$ )- CH  $_2$ -, where R  $^{12}$  is as defined above, or

12

carbon atoms and where  $\mathbb{R}^{13}$  is H or an alkyl group having alpha to the carbonyl with an alkyl group having 1 to 4 R<sup>1</sup> represents R<sup>13</sup>-NH-CO-alkyl-, where the alkyl group has 1 to 4 carbon atoms and is possibly substituted 1 to 4 carbon atoms or  $-CH_2COOR^{12}$ , where  $\mathbb{R}^{12}$  is as

20

defined above, or 25

having 1 to 4 carbon atoms and where  $\mathbb{R}^{12}$  is as defined substituted alpha to the carbonyl with an alkyl group  $\rm R^1$  represents  $\rm R^{12}00C\text{-}CH_2\text{--}00C\text{--}alkyl\text{--},}$  where the alkyl group has 1 to 4 carbon atoms and is possibly above, or

9

 $R^1$  represents  $R^{14} SO_2^{-}$ ,  $Ph(4-COOR^{12})-SO_2^{-}$ ,  $Ph(3-COOR^{12}) \mathrm{SO_2}^-$ ,  $\mathrm{Ph}(2\mathrm{-COOR}^{12})\mathrm{-SO_2}^-$  where  $\mathrm{R}^{12}$  is as defined above and R<sup>14</sup> is an alkyl group having 1-4 carbon atoms, or

35

 $R^{\rm l}$  represents -CO- $R^{\rm l5},$  wherein  $R^{\rm l5}$  is an alkyl group

having 1-4 carbon atoms, or

R<sup>1</sup> represents -CO-OR<sup>15</sup>, where R<sup>15</sup> is as defined above,

 ${ t R}^1$  represent -CO-(CH $_2$ ) $_{ t p}$ -COOR $^{12}$ , where  ${ t R}^{12}$  is as defined above and p is an interger 0, 1 or 2, or

 $\mathbb{R}^1$  represents -CH<sub>2</sub>PO(OR<sup>16</sup>)<sub>2</sub>, -CH<sub>2</sub>SO<sub>3</sub>H or each occurrence, H, methyl or ethyl;  $^{-CH}_2-(5-(1H)-\text{tetrazolyl})$  where  $\mathbb{R}^{16}$  is, individually at

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having 1 to 4 carbon atoms; carbon atoms and where  $\mathbb{R}^{21}$  is H or an alkyl group  $\mathtt{R}^2$  represents H or an alkyl group having 1 to 4 carbon atoms or  $R^{21}$ 00C-alkyl-, where the alkyl group has 1 to 4

15

fluorine atoms, or and the alkyl group may or may not carry one or more  ${f R}^3$  represents an alkyl group having 1-4 carbon atoms,

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group having 1 to 4 carbon atoms, or group which may or may not be substituted with an alkyl  $\mathbb{R}^3$  represents a cyclopentyl, cyclohexyl- or a phenyl

25

carbon atoms and k is 0, 1, or group, where  $\mathbb{R}^{21}$  is H or an alkyl group having 1 to 4  ${\tt R}^3$  represents a phenyl group substituted with a  ${\tt OR}^{31}$ 

 ${\mathtt R}^3$  represents a 1-naphthyl or 2-naphthyl group and k is 0, 1, or

 $\mathbf{R}^3$  represent a cis- or trans-decalin group and  $\mathbf{k}$  is 0,

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R<sup>3</sup> represents 4-pyridyl, 3-pyrrolidyl or 3-indolyl which may or may not be substituted with a  $0R^{31}$  group,

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where  $R^{31}$  is as defined above and k is 0, 1, or

cyclohexyl- or a phenyl group;  $\mathbb{R}^3$  represents  $\mathrm{Si}(\mathrm{Me})_3$  or  $\mathrm{CH}(\mathbb{R}^{32})_2$ , wherein  $\mathbb{R}^{32}$  is a

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atoms, a cyclohexyl or a phenyl group;  $R^4$  represents H, an alkyl group having 1 to 4 carbon

IIIb or IIIc  $\mathtt{A^2}$  represents a structural fragment of Formula IIIa,

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wherein:

20

25 p is an interger 0, 1 or 2;

m is an integer 1, 2, 3 or 4;

Y represents a methylene group, or

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atoms, a hydroxy group or an oxo group in position 4, membered ring may or may not carry one or two fluorine or may or may not be unsaturated, or Y represents an ethylene group and the resulting 5-

heteroatom functionality in position 4, or Y represents  $-CH_2-O-$ ,  $-CH_2-S-$ ,  $-CH_2-SO-$ , with the

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unsaturated in position 4 and 5, or carry in position 4 fluorine atom, a hydroxy group or an oxo group, carry Y represents a n-propylene group and the resulting 6membered ring may or may not carry in position 5 one two fluorine atoms in one of positions 4 or 5 or be an alkyl group with 1 to 4 carbon atoms, or

Y represents -CH<sub>2</sub>-0-CH<sub>2</sub>-, -CH<sub>2</sub>-S-CH<sub>2</sub>-, -CH<sub>2</sub>-SO-CH<sub>2</sub>-, or

Y represent -CH2-CH2-CH2-CH2-; ដ

R3 is as defined above;

 $R^5$  represents H or an alkyl group having 1 to 4 carbon atoms, or

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R<sup>51</sup> is H or an alkyl group having 1 to 4 carbon atoms;  $R^5$  represents  $-(CH_2)_p$ -COOR<sup>51</sup>, where p is 0, 1 or 2 and

n is an integer 0, 1, 2, 3 or 4; 20

B represents a structural fragment of Formula IVa, IVb, IVc or IVd

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wherein:

r is an interger 0 or 1;

 $\mathbf{X}^1$  represent  $\mathbf{CH}_2$  or  $\mathbf{NH}$  or is absent;

 $\chi^2$  represents  ${\rm CH}_2$ , NH or C=NH;

 $\chi^3$  represents NH, C=NH, N-C(NH)-NH<sub>2</sub>, CH-C(NH)-NH<sub>2</sub>, CH-NH-C(NH)-NH2 or CH-CH2-C(NH)-NH2; ព

X4 represents CH2 or NH;

 $\chi^5$  represents C(NH)-NH2 or NH-C(NH)-NH2;

X<sup>6</sup> represents CH or N;

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 $R^6$  is H or an alkyl group having 1-4 carbon atoms;

either the compound as such or stereoisomers thereof or in the form of a physiologically acceptable salt. 20

2. A compound of the general formula

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Formula V

30

wherein:

 ${\bf A}^1$  represents a structural fragment of Formula IIa, IIb, IIC, IId or IIe; 35

wherein:

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k is an integer 0, 1, 2, 3 or 4;

m is an integer 1, 2, 3 or 4;

q is an integer 0, 1, 2, or 3;

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carbon atoms, or a benzyl group, or 0, 1 or 2 and  $\mathbb{R}^{17}$  is,  $\mathsf{COOR}^{12}$ ,  $\mathsf{CONHR}^{12}$ , where  $\mathbb{R}^{12}$  is H, an alpha substituent is a group  $R^{17}$ -(CH<sub>2</sub>) $_p$ -, wherein p is position which is alpha to the carbonyl group, and the to 4 carbon atoms and is possibly substituted in the group, and  $\mathbb{R}^{11}$  is H or an alkyl group having 1 to 6 alkyl group having 1 to 4 carbon atoms or a benzyl  $R^1$  represents  $R^{11}00C$ -alkyl-, where the alkyl group has 1

30

 $\mathbb{R}^1$  represents Ph(4-COOR<sup>12</sup>)- CH<sub>2</sub>-, where  $\mathbb{R}^{12}$  is as defined above, or

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defined above, or 1 to 4 carbon atoms or  $-\mathrm{CH}_2\mathrm{COOR}^{12}$ , where  $\mathrm{R}^{12}$  is as carbon atoms and where R<sup>13</sup> is H or an alkyl group having alpha to the carbonyl with an alkyl group having 1 to 4 has 1 to 4 carbon atoms and is possibly substituted  $\mathtt{R}^1$  represents  $\mathtt{R}^{13}\mathtt{\_NH-CO-alkyl-}$ , where the alkyl group

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having 1 to 4 carbon atoms and where  $\mathbb{R}^{12}$  is as defined substituted alpha to the carbonyl with an alkyl group group has 1 to 4 carbon atoms and is possibly above, or  $\mathbb{R}^1$  represents  $\mathbb{R}^{12}$ 00C-CH $_2$ -00C-alkyl-, where the alkyl

10

15 so<sub>2</sub>-, and  $\mathbb{R}^{14}$  is an alkyl group having 1-4 carbon atoms, or  ${
m R}^1$  represents  ${
m R}^{14}{
m SO}_2$ -,  ${
m Ph}(4{
m -COOR}^{12}){
m -SO}_2$ -,  ${
m Ph}(3{
m -COOR}^{12}){
m -}$  $^{
m Ph}(2 ext{-COOR}^{12}) ext{-SO}_2 ext{-}$  where  $^{
m R}^{12}$  is as defined above

 $\mathbb{R}^1$  represents -CO- $\mathbb{R}^{15}$ , wherein  $\mathbb{R}^{15}$  is an alkyl group having 1-4 carbon atoms, or

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 $\mathbb{R}^1$  represents -CO-OR<sup>15</sup>, where  $\mathbb{R}^{15}$  is as defined above, 유

25 above and p is an interger 0, 1 or 2, or  $\mathbb{R}^1$  represent -CO-(CH<sub>2</sub>)<sub>p</sub>-COOR<sup>12</sup>, where  $\mathbb{R}^{12}$  is as defined

having 1 to 4 carbon atoms or a benzyl group; carbon atoms and where  $\mathbb{R}^{21}$  is H or an alkyl group atoms or  $R^{21}$ 00C-alkyl-, where the alkyl group has 1 to 4  ${f R}^2$  represents H or an alkyl group having 1 to 4 carbon

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fluorine atoms, or and the alkyl group may or may not carry one or more R<sup>3</sup> represents an alkyl group having 1-4 carbon atoms,

group which may or may not be substituted with an alkyl R<sup>3</sup> represents a cyclopentyl, cyclohexyl- or a phenyl

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group having 1 to 4 carbon atoms, or

group, where  $\mathbb{R}^{31}$  is H or an alkyl group having 1 to 4  $\mathtt{R}^3$  represents a phenyl group substituted with a  $\mathtt{OR}^{33}$ carbon atoms and k is 0, 1, or  $R^3$  represents a 1-naphthyl or 2-naphthyl group and k is 0, 1, or

 $R^3$  represent a cis- or trans-decalin group and k is  $\theta,$ 2

which may or may not be substituted with a  $OR^{31}$  group, R<sup>3</sup> represents 4-pyridyl, 3-pyrrolidyl or 3-indolyl where R31 is as defined above and k is 0, 1, or

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 $R^3$  represents Si(Me)  $_3$  or  ${\rm CH(R^{32})}_2$  , wherein  $R^{32}$  is a cyclohexyl- or a phenyl group;  $R^4$  represents H, an alkyl group having 1 to 4 carbon atoms, a cyclohexyl or a phenyl group; 2

 $\mathtt{A}^2$  represents a structural fragment of Formula IIIa, IIIb or IIIc

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wherein:

p is an interger 0, 1 or 2;

m is an integer 1, 2, 3 or 4;

Y represents a methylene group, or

membered ring may or may not carry one or two fluorine atoms, a hydroxy group or an oxo group in position 4, Y represents an ethylene group and the resulting 5or may or may not be unsaturated, or

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Y represents -CH2-0-, -CH2-S-, -CH2-S0-, with the heteroatom functionality in position 4, or

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unsaturated in position 4 and 5, or carry in position 4 Y represents a n-propylene group and the resulting 6fluorine atom, a hydroxy group or an oxo group, carry membered ring may or may not carry in position 5 one two fluorine atoms in one of positions 4 or 5 or be an alkyl group with 1 to 4 carbon atoms, or

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Y represents -CH<sub>2</sub>-0-CH<sub>2</sub>-, -CH<sub>2</sub>-S-CH<sub>2</sub>-, -CH<sub>2</sub>-SO-CH<sub>2</sub>-, or

Y represent -CH2-CH2-CH2-;

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R<sup>3</sup> is as defined above;

 $R^5$  represents H or an alkyl group having 1 to 4 carbon atoms, or ဗ္ဗ

R<sup>51</sup> is H or an alkyl group having 1 to 4 carbon atoms;  $R^5$  represents -(CH<sub>2</sub>)  $_{\rm p}$ -COOR<sup>51</sup>, where p is 0, 1 or 2 and

n is an integer 0, 1, 2, 3 or 4;

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B represents a structural fragment of Formula IVa, IVb,

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wherein:

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r is an interger 0 or 1;

X1 represent CH2 or NH or is absent;

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 $X^2$  represents  $CH_2$ , NH or C=NH;

NH-C(NH)-NH2 or CH-CH2-C(NH)-NH2;  $x^3$  represents NH, C=NH, N-C(NH)-NH<sub>2</sub>, CH-C(NH)-NH<sub>2</sub>, CH-

 $X^4$  represents  $CH_2$  or  $NH_7$ 

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X<sup>5</sup> represents C(NH)-NH<sub>2</sub> or NH-C(NH)-NH<sub>2</sub>;

X<sup>6</sup> represents CH or N;

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 $\mathbb{R}^6$  is H or an alkyl group having 1-4 carbon atoms;

D is 2 or (2)2;

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Z is a benzyloxy carbonyl group;

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in the form of a physiologically acceptable salt. either the compound as such or stereoisomers thereof or

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- a structural fragment of formula IIa or IIb. 3. A compound according to claims 1 or 2 wherein  $\mathbb{A}^1$  is
- atoms and R11 is H.  $\mathbb{R}^{11}$ 00C-alkyl-, where the alkyl group has 1 to 4 carbon precedings claims 1-3 wherein  $\mathbb{R}^1$  represents 4. A compound according to one or more of the
- precedings claims 1-4 wherein  $\mathbb{A}^2$  is a structural fragment of formula IIIa. 5. A compound according to one or more of the

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- formula IIIb. claims 1-4 wherein  ${\mathbb A}^2$  is a structural fragment of 6. A compound according to one or more of the preceding
- $C(NH)-NH_2$ , r is 1 and n is 1. claims 1-6 wherein B is a structural fragment of A compound according to one or more of the preceding formula IVa, wherein  $\chi^1$ ,  $\chi^2$  and  $\chi^4$  are  $CH_2$ ,  $\chi^3$  is CH-

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C(NH)-NH2, r is 0 or 1 and n is 1 or 2. formula IVa, wherein  $X^1$ ,  $X^2$  and  $X^4$  are  $CH_2$ ,  $X^3$  is Nclaims 1-6 wherein B is a structural fragment of 8. A compound according to one or more of the preceding

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claims 1-6 wherein B is a structural fragment of 9. A compound according to one or more of the preceding formula IVb, wherein  $X^5$  is C(NH)- $NH_2$  and  $R^6$  is H and n

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of formula IVa, wherein  $x^1$  and  $x^3$  are NH,  $x^2$  is C=NH,  $x^4$ preceding claims 1-6 wherein B is a structural element 10. A compound according to one or more of the

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is  $CH_2$ , r is 1 and n is 2.

of formula IVa, wherein  $x^1$  is absent,  $x^2$  and  $x^4$  are  $\mathrm{CH}_2$ , preceding claims 1-6 wherein B is a structural element 11. A compound according to one or more of the  $\chi^3$  is N-C(NH)-NH2, r is 0 and n is 1 or 2.

 $x^4$  are  $\text{CH}_2$  and  $x^3$  is  $\text{CH-C(NH)-NH}_2$  or  $\text{N-C(NH)-NH}_2,\ r$  is 0or 1, or  $x^1$  and  $x^3$  is NH,  $x^2$  is C=NH,  $x^4$  is CH2,  $\Gamma$  is 1, or  $\chi^1$  is absent,  $\chi^2$  and  $\chi^4$  are  $CH_2$ ,  $\chi^3$  is N-C(NH)-NH2, rn-propylene group and the resulting 6-membered ring may 12. A compound according to claims 1 or 2 in which n is wherein k is 0 or 1,  $R^1$  represents  $R^{11}00C$ -alkyl-, where structural fragment of formula IVa wherein  $\boldsymbol{x}^1,~\boldsymbol{x}^2$  and the alkyl group has 1 to 4 carbon atoms,  $\mathbb{R}^2$  represents represents a methylene group, an ethylene group, or a or may not carry in position 4 an alkyl group with 1 H,  $\mathrm{R}^3$  represents a cyclohexyl group,  $\mathrm{A}^2$ , represents a to 4 carbon atoms,  $\mathrm{R}^5$  represents H, B represents a 1 or 2,  $A^1$  is a structural fragment of formula IIa structural fragment of Formula IIIa wherein Y is o.

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13. A compound according to claims 1 or 2 in which n is 1,  $\mathbf{A}^1$  is a structural fragment of formula IIa wherein kn-propylene group and the resulting 6-membered ring may is 0 or 1,  $\mathrm{R}^1$  represents  $\mathrm{R}^{11}$ 00C-alkyl-, where the alkyl or may not carry in position 4 an alkyl group with 1 represents a methylene group, an ethylene group, or a to 4 carbon atoms,  $\mathrm{R}^5$  represents H, B represents a group has 1 to 4 carbon atoms,  $\mathrm{R}^2$  represents H,  $\mathrm{R}^3$ structural fragment of formula IVb wherein  $\mathrm{x}^5$ represents a cyclohexyl group,  $\lambda^2$  represents a structural fragment of Formula IIIa wherein Y represents  $C(NH)-NH_2$  and  $R^6$  is H.

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14. A compound selected from

HOOC-CH2-(RorS) CH(COOH)-(R) Cha-Aze-Pab/b HOOC-CH2-(ROYS) CH(COOH)-(R) Cha-Aze-Pab/a HOOC-CH2-(R, S) CH(COOH)-(R) C91-Pic-Pab HOOC-CH2-(R,S)CH(COOH)-(R)Cha-Aze-Pab HOOC-CH2-NH-CO-CH2-(R) Cha-Aze-Pab HOOC-CH2-CH2-(R) Cha-Aze-Pab HOOC-CH2-CH2-(R) Cgl-Pro-Pab HOOC-CH2-CH2-(R) Cgl-Aze-Pab (HOOC-CH2) 2-(R) Cg1-Pro-Pab HOOC-CH2-(R) Cha-Aze-Pab HOOC-CH2-(R) Cha-Pro-Pab HOOC-CH2-(R) Cgl-Pro-Pab HOOC-CH2-(R)Cgl-Aze-Pab H-(R)Cha-Aze-Pab H-(R) Cha-Pro-Pab H-(R)Cgl-Pic-Pab 12 ខ្ព

HOOC-CH2-(RorS) CH(COOH)-(R) Cha-Pic-Pab/a HOOC-CH2-(ROIS) CH(COOH)-(R) Cha-PIO-Pab/b HOOC-CH2-(RorS)CH(COOH)-(R)Cha-Pro-Pab/a HOOC-CH2-NH-CO-CH2-(R) Cha-Pro-Pab Etooc-CH2-CH2-CH2-(R) Cha-Pro-Pab HOOC-CH2-CH2-(Me) (R) Cha-Pro-Pab Ph (4-COOH) -SO2-(R) Cha-Pro-Pab HOOC-CH2-CH2-(R) Cha-Pro-Pab HOOC-CH2-(R) Cha-Pic-Pab H-(R)Cha-Pic-Pab 25

HOOC-CH2-(Me) (R) Cha-Pro-Pab

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HOOC-CH2-(Rors) CH(COOH)-(R) Cha-Pic-Pab/b Me-00C-CH2-CO-(R) Cha-Pic-Pab HOOC-CH2-CH2-(R) Cha-Pic-Pab HOOC-CH2-CO-(R) Cha-Pic-Pab H2N-CO-CH2-(R) Cha-Pic-Pab HOOC-CO-(R)Cha-Pic-Pab 35 30

Boc-(R) Cha-Pic-Pab Ac-(R)Cha-Pic-Pab

H-(R)Hoc-Aze-Pab H-(R)Cha-(R,S)betaPic-Pab HOOC-CH2-CH2-(R) Cha-Val-Pab HOOC-CH2-(R) Cha-Val-Pab HOOC-CH2-CH2-(R)Cha-(R,S)betaPic-Pab Me-SO<sub>2</sub>-(R)Cha-Pic-Pab

HOOC-CH2-(R)Hoc-Pic-Pab HOOC-CH2-(R,S)CH(COOH)-(R)Hoc-Pro-Pab (HOOC-CH<sub>2</sub>)<sub>2</sub>-(R)Hoc-Pic-Pab HOOC-CH2-CH2-(R) Hoc-Aze-Pab

H-(R)Cha-Aze-Pig HOOC-CH2-(R)Cgl-Pro-Pig HOOC-CH2-CH2-(R) Cgl-Aze-Pig HOOC-CH2-CH2-(R)Tic-Pro-Pab  $\mathtt{HOOC-CH_2-CH_2-(R)Pro(3-(S)Ph)-Pro-Pab}$ HOOC-CH<sub>2</sub>-(R)Pro(3-(S)Ph)-Pro-Pab

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MeOOC-CH<sub>2</sub>-(R)Cgl-Aze-Pab HOOC-(R,S)CH(Me)-(R)Cha-Pro-Pab H-(R)Cgl-Aze-Pab H-(R)Cgl-Ile-Pab

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H-(R)Cha-Pro-Pac HOOC-CH2-(R)Cgl-Aze-Pac

H-(R)Cgl-Pro-Pac "HexOOC-CH2-(R)Cgl-Aze-Pab <sup>n</sup>BuOOC-CH<sub>2</sub>-(R)Cgl-Aze-Pab Et00C-CH<sub>2</sub>-(R) Cg1-Aze-Pab

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HOOC-CH2-(R) Cha-Pro-Pac

H-(R)Cha-Pic-(R,S)Itp  $HOOC-CH_2-(R)$  Cha-Aze-(R, S) Itp HOOC-CH2-(R)Cgl-Aze-(R,S)Itp HOOC-CH2-CH2 (HOOC-CH2)-(R) Cha-Pro-Pig (HOOC-CH<sub>2</sub>)<sub>2</sub>-(R)Cgl-Pro-Pig HOOC-CH2-CH2-(R) Cha-Pro-Pig HOOC-CH2-(R)Cha-Pro-Pig HOOC-CH2-(R) Cha-Aze-Pig HOOC-CH2-CH2-(R) Cha-Aze-Pac HOOC-CH2-CH2-(R)Cgl-Pro-Pac •

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H-(R)Cha-Pro-(R,S)Nig HOOC-CH2-(R)Cha-Pro-(S)Itp HOOC-CH2-CH2-(R) Cha-Aze-Rig HOOC-CH2-(R) Cha-Pro-Rig HOOC-CH<sub>2</sub>-(R)Cgl-Aze-Rig H-(R)Cgl-Aze-Rig H-(R)Cha-Pro-(R,S)Hig HOOC-CH2-(R)Cgl-Pro-(R,S)Hig H-(R)Cgl-Pro-(R,S)Hig HOOC-CH2-(R)Cha-Pic-(R,S)Itp

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H-(R)Cha-Pro-Mig

15 of a physiologically acceptable salt. either as such or stereoisomer thereof or in the form H-(R)Cha-Aze-Dig H-(R)Cha-Pro-Dig

15. A compound selected from

20 HOOC-CH2-(R) Cha-Pro-Pig HOOC-CH2-(R) Cha-Pro-Pac EtOOC-CH2-(R)Cgl-Aze-Pab HOOC-CH2-(R)Cgl-Pro-Pig HOOC-CH2-(R)Cha-Pic-Pab HOOC-CH2-(R) Cha-Pro-Pab HOOC-CH2-CH2-(R) Cha-Aze-Pab HOOC-CH2-CH2-(R)Cha-Pro-Pab HOOC-CH<sub>2</sub>-(R)Cgl-Aze-Pab

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မ of a physiologically acceptable salt. either as such or stereoisomer thereof or in the form

16. A compound selected from

 $BnOOC-CH_2-(R)Cgl-Pro-Pab(Z)$  $BnOOC-CH_2-CH_2-(R)Cgl-Aze-Pab(Z)$  $Bn00C-CH_2-(R)Cgl-Aze-Pab(Z)$ 

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Bnooc-(R, S) CH(Me) -(R) Cha-Pro-Pab(Z)

4eooc-CH2-(R) Cg1-Aze-Pab(Z)

 $^{n}$ Buooc-CH<sub>2</sub>-(R)Cgl-Aze-Pab(Z)

 $^{n}$ HexOOC-CH<sub>2</sub>-(R)Cg1-Aze-Pab(Z)

BnOOC-CH<sub>2</sub>-(RorS)CH(COOBn)-(R)Cha-Aze-Pab(Z)/a Bnooc-CH<sub>2</sub>-(RorS)CH(COOBn)-(R)Cha-Aze-Pab(Z)/b

Bnooc-CH2-NH-CO-CH2-(R) Cha-Aze-Pab(Z)

Bnooc-CH2-CH2-(R) Cha-Aze-Pab(Z)

Bnooc-CH<sub>2</sub>-(R,S)CH(COOBn)-(R)Cha-Aze-Pab(Z)

Bnooc-CH2-(R) Cha-Aze-Pab(Z)

Bnooc-CH2-(R,S)CH(COOBN)-(R)Cgl-Pic-Pab(Z)

 $\mathtt{Bnooc-CH_2-CH_2-(R)\,Cgl-Pro-Pab(Z)}$ 

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(Bnooc-CH<sub>2</sub>)<sub>2</sub>-(R)Cgl-Pro-Pab(Z)

Bnooc-CH2-CH2-(R) Cha-Aze-Pac(Z)

 $Bnooc-CH_2-(R)Cha-Aze-Pig(Z)$ 

Bnooc-CH2-CH2-(R) Cha-Pro-Pig(Z)

Bnooc-CH2-CH2 (Bnooc-CH2)-(R) Cha-Pro-Pig(Z)

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Bn00C-CH2-(R, S) CH(C00Bn)-(R) Cha-Pro-Pab(Z)

Bnooc-CH2-CH2-(Me) (R) Cha-Pro-Pab(Z)

Bnooc-CH2-(Me) (R) Cha-Pro-Pab(Z) Bnooc-CH2-CH2-(R) Cha-Pro-Pab(Z)

Bnooc-CH2-(R) Cha-Pro-Pab(Z)

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Bn00C-CH2-NH-CO-CH2-(R)Cha-Pro-Pab(Z)

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Ph (4-COOH) -SO2-(R) Cha-Pro-Pab(Z)

Bnooc-CH2-(R) Cha-Pro-Rig(Z)

either as such or stereoisomer thereof or in the form

17. A compound selected from

 $Bnooc-cH_2-(R) Cgl-Aze-Pab(Z)$ 

Bn00C-CH2-(R)Cha-Pro-Pab(Z) Bnooc-CH2-(R) Cha-Pic-Pab(Z)

Bnooc-cH2-(R)cgl-Pro-Pig(Z)2

 $Bnooc-CH_2-(R)Cha-Pro-Pac(Z)$  $Etcoc-CH_2-(R)Cgl-Aze-Pab(Z)$ 

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BnOOC-CH2-(R, S) CH(COOBn)-(R) Hoc-Pro-Pab(Z)

Snooc-CH2-CH2-(R) Cha-(R, S) Val-Pab(Z)

Snooc-CH2-(R)Cha-Val-Pab(Z)

4e-SO2-(R) Cha-Pic-Pab(Z)

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Ac-(R) Cha-Pic-Pab(Z)

Bnooc-CH2-CH2-(R) Hoc-Aze-Pab(Z)

Bn00C-CH2-CH2-(R) Pro(3-(S) Ph) -Pro-Pab(Z) Bnooc-CH2-(R) Pro(3-(S) Ph) -Pro-Pab(Z)

Bnooc-CH2)2-(R)Hoc-Pic-Pab(Z)

Bnooc-CH2-(R) Hoc-Pic-Pab(Z)

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 ${\tt Bnooc-CH}_2{\tt -CH}_2{\tt -(R)\,Cgl-Aze-Pig(Z)}_2$ 

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Bnooc-CH2-(R)Cgl-Pro-Pig(Z)2

Bnooc-CH2-(R) Cgl-Aze-Pac(Z)

Bnooc-CH2-CH2-(R)Tic-Pro-Pab(Z)

Bnooc-CH2-(R) Cha-Pro-Pig(Z)

either as such or stereoisomer thereof or in the form of a physiologically acceptable salt.

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 $Etooc-cH_2-(R)cgl-Aze-Pab(Z)$ 

 ${\tt Bnooc-CH_2-CH_2-(R)\,Cgl-Pro-Pac(Z)}$  $Bnooc-CH_2-(R)Cha-Pro-Pac(Z)$ 

 $Bnooc-cH_2-(R) Cha-Pro-Pig(Z)$ 

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(Bnooc-CH2)2-(R)Cgl-Pro-Pig(Z)

Shooc-CH2-(R) Cha-Pic-(R,S) Itp(Z)

3n00C-CH2-(R)Cgl-Pro-(R,S)Hig(Z) snooc-CH2-(R) Cg1-Aze-Rig(Z)

BnOOC-CH2-CH2-(R) Cha-Aze-Rig(Z)

of a physiologically acceptable salt.

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Bnooc-CH2-(R,S)CH(COOBn)-(R)Cha-Pic-Pab(Z)

Bnooc-CH<sub>2</sub>-(R) Cha-Pic-Pab(Z)

Boc-(R) Cha-Pic-Pab(Z)

Bnooc-CH<sub>2</sub>-CH<sub>2</sub>-(R)Cha-Pic-Pab(Z)

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Etooc-co-(R) Cha-Pic-Pab(Z)

Meooc-CH2-CO-(R) Cha-Pic-Pab(Z)

12N-CO-CH2-(R) Cha-Pic-Pab(Z)

18. A compound selected from

H-(R) Pro-Phe-Pab
HOOC-CH<sub>2</sub>-(R) Pro-Phe-Pab
H-(R) Phe-Phe-Pab
HOOC-CH<sub>2</sub>-(R) Phe-Phe-Pab
HOOC-CH<sub>2</sub>-(R) Phe-Phe-Pab

either as such or stereoisomer thereof or in the form of a physiologically acceptable salt.

19. A compound selected from

Boc-(R) Pro-Phe-Pab(Z)
BnOC-CH<sub>2</sub>-(R) Pro-Phe-Pab(Z)
Boc-(R) Phe-Phe-Pab(Z)
MeOOC-CO-(R) Phe-Phe-Pab(Z)
BnOOC-CH<sub>2</sub>-(R) Phe-Phe-Pab(Z)

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either as such or steroisomer thereof or in the forms of a physiologically accetable salt.

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20. A process for preparing a compound according to any of claims 1-19, which process comprises coupling of an N-terminally protected amino acid or dipeptide or amino acid, when an N-terminally amino acid is used a second aminoacid is added afterwards using standard methods to a compound

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2N (CH2), ---

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wherein n is an integer 0, 1, 2, 3 or 4, X is B or B-D, where B is as defined in formula I and D is as defined in formula V as such or having the guanidino or amidino nitrogens either mono or diprotected with an amine

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protecting group such as a benzyloxy carbonyl- or tertbutyloxy carbonyl- or p-toluenesulphonyl group or x is a group transferable into B followed by removal of the protecting group(s) or deprotection of the N-terminal nitrogen followed by alkylation of the N-terminal nitrogen and if desired deprotection by known methods and if desired forming a physiologically acceptable salt, and in those cases where the reaction results in a mixture of stereoisomers, these are optionally separated by standard chromatographic or re-crystallisation techniques, and if desired a single stereoisomer is isolated.

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21. A process according to claim 20 for preparing a compound according to any of claims 1-19, which process comprises:

a) (Method Ia) Coupling of an N-terminally protected dipeptide, selected from A<sup>1</sup> and A<sup>2</sup> in Formulas I or V by using standard peptide coupling, shown in the formula

 $w^1 - A^1 - A^2 - 0$ 

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 $H_2N$ — $(CH_2)$ n— $Q^1$ 

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 $w'-A^1-A^2-W-(CH_0)$ 

wherein n is as defined in Formula I,  $W^1$  is an N-teminal amino protecting group such as tert-butyloxy carbonyl and benzyloxy carbonyl and and  $Q^1$  is -C(NH)-

NH<sub>2</sub>, -C(NW<sup>2</sup>)-NH-W<sup>2</sup>, -C(NH)-NH-W<sup>2</sup>, -NH-C(NH<sup>2</sup>)-NH-W<sup>2</sup>, where W<sup>2</sup> is an amine protecting group such as tert-butyloxy carbonyl or benzyloxy carbonyl, or Q<sup>1</sup> is -CN, -CO-NH<sub>2</sub> or -CS-NH<sub>2</sub>, where the group is subsequently transferred into a amidino group or Q<sup>1</sup> is NH<sub>2</sub> or NH-W<sup>2</sup>, where W<sup>2</sup> is as defined above, where the amino group is subsequently transferred into a guanidino group (giving Q<sup>1</sup> = -NH-C(NH)-NH<sub>2</sub>), after deprotection of the W<sup>2</sup>-group when Q<sup>1</sup> is -NH-W<sup>2</sup> (W<sup>2</sup> in this case must be orthogonal to W<sup>1</sup>), by methods known in the art.

b) (Method 1b) Coupling of an N-terminally protected amino acid, selected from  $\mathbb{A}^2$  in Formulas I or V and prepared by using standard peptide coupling, shown in the formula

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$$w^{\prime}$$
—  $A^2$  — $a_1$ 

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$$\begin{vmatrix}
H_2N-(CH_2)_n \\
W'-A^2 & W'-CH_2 \\
W'-A^2 & W'-CH_2 \\
W'-A^2 & W'-CH_2 \\
W'-CH_2 & W'-CH_2 \\
W'-CH_2$$

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wherein n,  $W^1$ , and  $Q^1$  are as defined above followed by deprotection of the  $W^1$ -group and coupling with the N-terminal amino acid, in a protected form, leading to the protected peptide described in Method Ia above,

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c) (Method IIa) Coupling of an N-terminally protected dipeptide, selected from  ${\bf A}^1$  and  ${\bf A}^2$  in Formulas I or V by using standard peptide coupling, shown in the formula

$$w'-A^1-A^2-\alpha$$

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$$^{\text{w}}$$
  $\mathsf{A}^1$   $\mathsf{A}^2$   $^{\text{hN}}$   $^{\text{cH}_2)}$   $^{\text{n}}$ 

wherein n is as defined in Formula I,  $W^1$  is an N-teminal amino protecting group such as tert. butyloxy carbonyl and and  $Q^1$  is -C(NH)-NH- $W^2$ , -C(NH)-NH- $W^2$ , -C(NH)-NH- $W^2$ , -C(NH)-NH- $W^2$ ,  $-N(W^2)$ -NH- $W^2$ ,  $-N(W^2)$ -NH- $W^2$ , where  $W^2$  is an amine protecting group such as tert. butyloxy carbonyl or benzyloxy carbonyl, or  $Q^1$  is -C(W)-CO-NH<sub>2</sub> or CS-NH<sub>2</sub>, where the group is subsequently transferred into a amidino group or  $Q^1$  is  $NH_2$  or NH- $W^2$ , where  $W^2$  is as defined above, where the amino group is subsequently transferred into a quanidino group  $Q^1$  is -NH-C(NH)-NH- $Q^2$ ), after deprotection of the  $W^2$ -group when  $Q^1$  is  $-NH-W^2$  ( $W^2$  in this case must be orthogonal to  $W^2$ ), by methods known in the art.

d) (Method IIb) Coupling of an N-terminally protected amino acid, selected from  ${\bf A}^2$  in Formulas I or V by using standard peptide coupling, shown in the formula

$$\begin{array}{c}
w' - A^2 - GH \\
\downarrow H_2N - (GH_2) - GH \\
\downarrow W' - A^2 - HN - (GH_2) - GH
\end{array}$$

wherein n,  $W^1$  and  $Q^1$  are as defined above followed by deprotection of the  $W^1$ -group and coupling with the N-terminal amino acid, in a protected form, leading to the protected peptide described in Method IIa above,

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e) (Metod IIIa) Coupling of an N-terminally protected dipeptide, selected from  $\mathbb{A}^1$  and  $\mathbb{A}^2$  in Formulas I or V by using standard peptide coupling, shown in the formula

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$$W' - A' - A^{-} - OH$$

$$H_{2}N - (CH_{2})_{1} - X_{1}^{*} - X_{2}^{*}$$

$$X' - X^{2} - X_{2}^{*} - X_{3}^{*} - X_{4}^{*}$$

$$W' - A^{1} - A^{2} - NH - (CH_{2})_{1} - X_{4}^{*}$$

$$W' - A^{1} - A^{2} - NH - (CH_{2})_{1} - X_{4}^{*}$$

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wherein n is as defined in Formula I and r is 0 or 1 when x², x² and x⁴ are CH₂ or r is 0 when x² and x⁴ are CH₂ or r is 0 when x² and x⁴ are protecting group such as tert-butyloxy carbonyl and benzyloxy carbonyl and and Q² is -C(NH)-NH₂, -C(NW²)-NH₂ or -C(NH)-NH-w², where w² is an amine protecting group such as tert-butyloxy carbonyl or benzyloxy carbonyl, or Q² is equal to w² where the amino group, after deprotection of the w² group (w² in this case must guanidino group using a unprotected, N-protected or in the art,

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f) (Method IIIb) coupling of an N-terminally protected amino acid, selected from  $\mathbb{A}^2$  in Formulas I or V by using standard peptide coupling, shown in the formula

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wherein n, r,  $x^1$ ,  $x^2$  and  $x^4$ ,  $w^1$ , and  $q^2$  are as defined above followed by deprotection of the  $w^1$ -group and coupling with the w-terminal amino acid, in a protected form, leading to the protected peptide described in Method IIIa above,

dipeptide, selected from  ${\bf A}^1$  and  ${\bf A}^2$  in Formulas I or V by Gethod IVa) Coupling of an N-terminally protected using standard peptide coupling, shown in the formula

$$w'-A^1-A^2-\alpha$$

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$$^{\text{w'}}-\mathbf{A}^1-\mathbf{A}^2-^{\text{MH}-(CH_2)_n}$$

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carbonyl or benzyloxy carbonyl and  $\ensuremath{\text{W}}^3$  is H or an amino terminal amino protecting group such as tert-butyloxy protecting group such as aryl sulfonyl, benzyloxy wherein n is as defined in Formula I,  $W^1$  is an Ncarbonyl or tert-butyloxy carbonyl.

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h) (Method IVb) Coupling of an N-terminally protected using standard peptide coupling, shown in the formula amino acid, selected from  ${\tt A}^2$  in Formulas I or V by

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$$w'$$
—  $A^2$  — $\alpha$ 

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$$\mathbf{w}^{1} - \mathbf{A}^{2} \xrightarrow{\mathbf{N}^{4} - (\mathbf{C}H_{2})_{n}} \mathbf{A}^{4}$$

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N-terminal amino acid, in a protected form, leading to wherein n,  $W^1$ , and  $W^3$  are as defined above followed by deprotection of the  $W^1$ -group and coupling with the the protected peptide described in Method IVa, the final compounds can be made in any of the following  $C(NH)-NH-W^2$ ,  $-N(W^2)-C(NH)-NH-W^2$  or  $-NH-C(NW^2)-NH-W^2$  ( $W^2$  $NH_2$ ,  $-NH-C(NH)-NH-W^2$ ,  $-N(W^2)-C(NH)-NH-W^2$  or  $-NH-C(NW^2)-C(NH)$ ways, depending on the nature of the  $\mathbb{Q}^1-$  or  $\mathbb{Q}^2-$ groups alkylation of the N-terminal nitrogen and if desired  $NH-W^2$ ), or a selective deprotection of the  $W^1-$  group (e.g when  $Q^1$  or  $Q^2 = -C(NW^2) - NH - W^2$ ,  $-C(NH) - NH - W^2$ , -NH used: Removal of the protecting group(s) (when  $\mathbf{Q}^{1_{\mathrm{cl}}}$  in this case must be orthogonal to  $W^1$ ) followed by C(NH)-NH<sub>2</sub>, -C(NW<sup>2</sup>)-NH-W<sup>2</sup>, -C(NH)-NH-W<sup>2</sup>, -NH-C(NH)deprotection.

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22. A compound according to any of claims 1-19 for use in therapy.

23. A compound according to any of claims 1-5 or 7-17 for use as an anticoagulant or antithrombotic agent.

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24. A compound according to any of claims 1-4, 6-10 or 18-19 for use as an antiinflammatory agent.

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effective amount of a compound as outlined in claims 1-19 in conjunction with one or more pharmaceutical 25. A pharmaceutical preparation comprising an carriers.

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pharmaceutical carriers for use as an anticoagulant or effective amount of a compound as outlined in any of claims 1-5 or 7-17 in conjuction with one or more 26. A pharmaceutical preparation comprising an antithrombotic agent.

27. A pharmaceutical preparation comprising an effective amount of a compound as outlined in any of claims 1-4, 6-10 or 18-19 in conjuction with one or more pharmaceutical carriers for use as an antiinflammatory agent.

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28. Use of compound according to any of claims 1-5 or 7-17 as an active ingredient for manufacture of a pharmaceutical preparation for inhibition of thrombin in a human or animal organism.

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29. Use of compound according to any of claims 1-4, 6-10 or 18-19 as an active ingredient for manufacture of a pharmaceutical preparation for inhibition of kininogenases in a human or animal organism.

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30. A method for obtaining inhibition of thrombin in a human or animal organism in need of such inhibition, comprising administering to said organism an inhibitory effective amount of a compound claimed in any of claims 1-5 or 7-17.

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31. A method for obtaining inhibition of kininogenases in a human or animal organism in need of such inhibition, comprising administering to said organism an inhibitory effective amount of a compound claimed in any of claims 1-4, 6-10 or 18-19.

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32. Use of a compound of the formula:

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either as such or having the amidino group either mono- or diprotected at the nitrogens with a protective group, or in the form of a salt, as a starting material in synthesis of a peptidic serine protease inhibitor, and in particular in synthesis of a peptidic thrombin inhibitor or a peptidic kininogenases inhibitor.

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33. A structural fragment of the formula

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as a structural element in a pharmaceutically active compound, especially a peptidic compound.

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34. A compound 4-aminomethyl-1-(N-benzyloxycarbonylamidino) benzene either as such, in the form of a salt or having a protection with a benzyloxycarbonyl group at the other nitrogen.

35. Use of a compound of the formula:

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either as such or having the amidino group either monoor diprotected at the nitrogens with a protective group, or in the form of a salt, as a starting material in synthesis of a thrombin inhibitor, and in particular in synthesis of a peptidic thrombin inhibitor.

36. A structural fragment of the formula

as a structural element in a thrombin inhibitor, especially a peptidic thrombin inhihitor.

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37. A compound 4-aminomethyl-1-(N-benzyloxy-carbonylamidino) cyclohexane either as such, in the form of a salt or having a protection with a benzyloxycarbonyl group at the other nitrogen.

38. A compound of the formula:

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either as such or having the amidino group either monoor diprotected at the nitrogens with a protective group, or in the form of a salt, .

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39. Use of a compound as described in claim 38, as a starting material in the synthesis of a serine protease inhibitor.

40. A structural fragment of the formula

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as a structural element in a thrombin inhibitor, especially a peptidic thrombin inhihitor.

41. A compound of the formula:

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either as such or having the amidino group either monoor diprotected at the nitrogens with a protective group, or in the form of a salt.

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42. Use of a compound as described in claim 41, as a starting material in the synthesis of a serine protease inhibitor.

43. A structural fragment of the formula

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as a structural element in a thrombin inhibitor, especially a peptidic thrombin inhihitor.

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44. A compound 4-aminoethyl-1-benzyloxy-carbonylamidino piperidine either as such, in the form of a salt or having a protection with a benzyloxycarbonyl group at the other nitrogen.

45. A compound of the formula:

HN CH2)

where n is 1 or 2 and s is 0 or 1, either as such or having the amidino group either monoor diprotected at the nitrogens with a protective group, or in the form of a salt.

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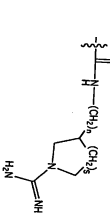
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46. Use of a compound as described in claim 45, as a starting material in the synthesis of a serine protease inhibitor.

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47. A structural fragment of the formula

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where n is 1 or 2 and s is 0 or 1, as a structural element in a thrombin inhibitor, especially a peptidic thrombin inhihitor.

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48. A compound which is (3RS)-1-(N-benzyloxycarbonyl-amidino)-3-aminomethyl pyrrolidine, (3RS)-1-(N-benzyl-oxycarbonylamidino)-3-aminoethyl pyrrolidine, 3-aminomethyl-1-(N-benzyloxycarbonyl-amidino) azetidine, 3-aminoethyl-1-(N-benzyloxycarbonyl-amidino) azetidine or either as such, in the form of a salt or having a protection with a benzyloxycarbonyl group at the other nitrogen.

INTERNATIONAL SEARCH REPORT

International application No. PCT/SE 94/00535

IPC © CO7K 5/06, CO7K 5/02, AG1K 38/55, CO7C 257/18, CO7C 257/16, CO7D 239/14, CO7D 211/26, CO7D 205/04, CO7O 207/09 describing to international Patent Charaffeation (IPC) or to both national described on and IPC Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched electronic data base consulted during the international search (name of data base and, where practicable, search terms used) Minimum documentation searched (classification system followed by classification symbols) A. CLASSIFICATION OF SUBJECT MATTER SE, DK, FI, NO classes as above B. FIELDS SEARCHED IPC6: A61K, C07K

BIOSIS, MEDLINE, EMBASE, CA, WPI, CLAIMS C. DOCUMENTS CONSIDERED TO BE RELEVANT

Relevant to claim No. 1-29 1-29 Caugory | Clation of document, with indication, where appropriate, of the relevant passages EP, A1, 0530167 (AKTIEBOLAGET ASTRA), 3 March 1993 (03.03.93)

EP, A1, 0513543 (BEHRINGMENKE AKTIENGESELLSCHAFT), 19 November 1992 (19.11.92)

EP, A1, 0236163 (SANOFI), 9 Sept 1987 (09.09.87)

1-29

EP, A2, 0542525 (ELI LILLY AND COMPANY), 19 May 1993 (19.05.93)

1-29

X See patent family annex. Further documents are listed in the continuation of Box C.

The later document published after the international filling date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the laveration document of particular relevance the claimed invention cannot be considered novel or examit be considered to invent we inventive step when the document is taken alone document defining the general state of the art which is not considered to be of particular relevance eries document but published on or after the international filling date eries document but published on or after the international filling date document which may throw doubts on priority claim(s) or which is cited to exabilish the publication date of mother claims or other special reason (as specified) Special categories of cited documents <u>`</u> ب ب

"y" document of particular relevance the claimed invention cannot be considered to inverve an inventive step when the document is combined with once or more other such documents, such combination being obvious to a person stilled in the Art. "&" document member of the same patent family document published prior to the international filing date but later than the priority date claimed document referring to an oral discionure, wer, exhibition or other means Ģ

Date of mailing of the international search report 31 -10- 1994 Authorized officer Date of the actual completion of the international search As to support AB2T
Name and mailing address of the ISA/
Swedish Patent Office
Box 5055, S-102 42 STOCKHOLM
Facsimile No. + 46 8 666 02 86
From PCT/ISA210 (second sheet) (July 1992) 27 October 1994

Elisabeth Carlborg Telephone No. +46 8 782 25 00

INTERNATIONAL SEARCH REPORT

International application No. PCT/SE 94/00535

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Relevant to claim No.	32,34	32,34	32,34	132,34	35,37
Clation of document, with indication, where appropriate, of the relevant passages	THROMBOSIS ET DIATHESIS HAEMORRHAGICA, Volume 23, No 3, 1970, j.D. Geratz, "Inhibition of Thrombin, Plasmin and Plasminogen Activation by Amidino Compounds" page 486 - page 499	THROMBOSIS ET DIATHESIS HAEMORRHAGICA, Volume 31, 1974, Erika Glusa et al, "The Influence of Benzamidine Derivatives on Human Platelet Function" page 172 - page 178	BIOCHEMICAL PHARMACOLOGY, Volume 23, 1974, Fritz Markwardt et al, "Synthetic Low Molecular Weight Inhibitors of Serum Kallikrein" page 2247 - page 2256	PHARMAZIE, Volume 34, No 10, 1979, D. Labes et al, "Hansch-Analyse der Hemmwirkung von 3- und 4- substitulerten Benzamidinen gegenüber Thrombin, Plasmin und Trypsin <sup>n</sup> page 649 - page 653	EP, A1, 0001774 (BAYER AKTIENGESELLSCHAFT ZENTRALBEREICH PATENTE), 16 May 1979 (16.05.79), page 32, line 16 - page 33, line 14; page 51, line 1 - line 9
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	寸	Clauson of document, with indication, where appropriate, of the relevant possesses THROMBOSIS ET DIATHESIS HAEMORRHAGICA, Volume 23, No 3, 1970, j.D. Geratz, "Inhibition of Thrombin, Plasmin and Plasminogen Activation by Amidino Compounds" page 486 - page 499	Claudon of document, with indication, where appropriate, of the relevant postages THROMBOSIS ET DIATHESIS HAEMORRHAGICA, Volume 23, No 3, 1970, j.D. Geratz, "Inhibition of Thrombin, Plasmin and Plasminogen Activation by Amidino Compounds" page 486 - page 499  THROMBOSIS ET DIATHESIS HAEMORRHAGICA, Volume 31, 1974, Erika Glusa et al, "The Influence of Benzamidine Derivatives on Human Platelet Function" page 172 - page 178	Claudon of document, with indication, where appropriate, of the relevant passages THROWBOSIS ET DIATHESIS HAEWORRHAGICA, Volume 23, No 3, 1970, j.D. Geratz, "Inhibition of Thrombin, plasmin and Plasminogen Activation by Amidino Compounds" page 486 - page 499  THROWBOSIS ET DIATHESIS HAEWORRHAGICA, Volume 31, 1974, Erika Glusa et al, "The Influence of Benzamidine Derivatives on Human Platelet Function" page 172 - page 178  BIOCHEMICAL PHARMACOLOGY, Volume 23, 1974, Fritz Markwardt et al, "Symthetic Low Molecular Weight Inhibitors of Serum Kallikrein" page 2247 - page 2256	Claudon of document, with indication, where appropriate, of the relevant passage THROMBOSIS ET DIATHESIS HAEMORRHAGICA, Volume 23, No. 3, 1970, j.D. Geratz, "Inhibition of Thrombin, plasmin and Plasminogen Activation by Amidino Compounds" page 486 - page 499  THROMBOSIS ET DIATHESIS HAEMORRHAGICA, Volume 31, 1974, Erika Glusa et al, "The Influence of Benzamidine Derivatives on Human Platelet Function" page 172 - page 178  Fritz Markwardt et al, "Synthetic Low Molecular Weight Inhibitors of Serum Kallikrein" page 2247 - page 2256  PHARMAZIE, Volume 34, No 10, 1979, D. Labes et al, "Hansch-Analyse der Hemmwirkung von 3- und 4-substitulerten Benzamidinen gegenüber Thrombin, Plasmin und Trypsin" page 649 - page 653

## INTERNATIONAL SEARCH REPORT

	pplicant, this international search report sently, this international search report is Nos.:	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Noz.:  No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nog.:  The additional search fees were accompanied by the applicant's protest.  No protest  No protest No protest secompanied the payment of additional search fees.
	fee, this Authority did not invite payment	1 🗀
	is international search report covers the	* See extre sheet  1. X As all required additional search fees were timely paid by the applicant, this international search report causes
	A. C.	a) claims 1-29, 33, 36, 40, 43 and 47, b) claims 32, 34, 35 and 37, c) claims 38 and 39, d) claims 41, 42, 44-46 and 48
	of first sheet)	Dis International Searching Authority found multiple inventions in this international spoileation, as follower.
<del>-</del>	second and third sentences of Rule 6.4(s	, —
<u> </u>	ly with the prescribed requirements to su भ्रोपुरः	2. X Claims Nos.: 33, 36, 40, 43 and 47 because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no mentingful international search can be carried out, specifically:  ** See Sature sheet
	ertain claims under Article 17(2)(a) for the following reason reched by this Authority, namely: treetment of the human or animal as diagnostic methods.	This international search report has not been established in respect of certain claims under Article 17(2)(s) for the following reasons:  1.     Chima Now: 30 and 31   Chima Now: 30 a
$\perp$	lon of Item 1 of first sheet)	Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
	PCT/SE 94/00535	1

## INTERNATIONAL SEARCH REPORT

International application No.
PCT/SE 94/00535

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The subjects of invention as listed below are so different from each other, that no technical relationship can be appreciated to be present so as to form a single general inventive concept.

Claims 1-29, 33, 36, 40, 43 and 47 relate to compounds, to processes for preparing them, to pharmaceutical preparations containing the compounds, to the use of the compounds for manufacture of pharmaceutical preparations and to structural fragments, which have been regarded as being part of the compounds disclosed in claim 1.

Claims 32, 34; 35 and 37 relate to structurally similar compounds (intermediates) and to the use of such compounds as starting material in synthesis of enzyme inhibitors.

Claims 38 and 39 relate to a compound (intermediate) and to its use as starting material in the synthesis of a serine protease inhibitor.

Claims 41, 42, 44-46 and 48 relate to compounds (intermediates) and to their use as starting material in the synthesis of a serine protease inhibitor.

\*\* The wording of claims 33, 36, 40, 43 and 47, which indicates several possible end compounds with a structural fragment, does not define one solution of one technical problem. Therefore, the claims are not searchable.

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International application Ive. PCT/SE 94/00535	Publication date	10/02/94 05/04/93 18/03/93 13/07/94 28/02/94 00/00/00 27/06/94 18/03/93	14/07/94 12/11/92 23/12/92 12/11/92 22/06/93 27/06/94	07/08/87 06/10/87 11/12/90 09/09/87 07/08/87	03/03/94 13/05/93 13/05/93 21/07/93 28/07/94 31/08/93 12/10/94	03/05/79 13/06/79
	Patent family member(s)	313 2499092 2116527 0605462 940945 9400589 243675 9305069	651196 1608692 1067249 4115468 5155898 242668	2593812 62228050 4977168 0236164 2593814 4791102	646767 2829892 2831292 2082748 1074438 925124 65858 5221964 525566	2748295 54073702
n 01/10/94	Paten	AP-A- AU-A- CA-A- EP-A- FI-A,D- HU-D- NO-A-	AU-B- AU-A- CN-A- DE-A- NZ-A-	. R-A,B- JP-A- US-A- EP-A,B- RR-A,B- US-A-	AU-B- AU-A- CA-A- CN-A- CN-A- UP-A- US-A- US-A-	DE-A- JP-A-
RNATIONAL SEARCH REFY	Publication date	03/03/93	19/11/92	09/09/87	19/05/93	16/05/79
INTERNATIONAL SEARCH NEFORT information on patent family members	Patent document cited in search report	0530167	0513543	0236163	0542525	0001774
EN	Patent d	EP-A1-	EP-A1-	EP-41-	EP-A2-	EP-A1-

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